Label-free detection: technologies, key considerations, and applications

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Why use label-free detection for binding interaction analysis?

Interactions between biomolecules serve as key triggers for many biological processes and, therefore, provide perfect targets for drug discoveries. Biological binding interactions are a dynamic process driven by changes to the environment. Therefore, techniques used to characterize these interactions needs to mirror the level of biological complexity in order to fully understand these systems.

LABEL-FREE BIOSENSOR BINDING ASSAY METHODS

Commonly used label-free binding analysis platforms include Bio-Layer Interferometry (BLI), Surface Plasmon Resonance (SPR), Isothermal Titration Calorimetry (ITC) and Microscale Thermophoresis (MST). In this e-book, we’ll compare the four technologies in terms of their capabilities and workflows, and provide guidance for choosing the most suitable platform based on assay needs and application.

BLI and SPR monitor binding interactions based on molecular accumulation that take place during complex formation. The binding complex is established at the biosensor surface by immobilization of one binding partner (ligand) and directly monitoring the binding of the analyte supplied from solution. The complex formation and dissociation are monitored in real time, providing kinetics and affinity data.

Bio-Layer Interferometry

BLI is an optical technology that measures the changes in interference pattern between light waves. ForteBio’s BLI-based platforms measure light interference originating from the tip of the biosensor surface where light wavelengths are made to reflect from two layers: a biocompatible layer at the end of the biosensor surface and an internal reference layer (Figure 1). Incident white light reflecting from the two layers interfere constructively or destructively depending on the thickness of the molecular biolayer at the biosensor tip. The spectral pattern of the reflected light changes as a function of the optical thickness of the molecular layer, i.e. the number of molecules bound to the biosensor surface. This spectral shift is monitored at the detector, and reported on a sensorgram as a change in wavelength (nm shift). Monitoring the interference pattern (i.e. spectral shift) in real time provides kinetics data on molecular interactions.

Dip and Read™ biosensors are core to BLI technology. The biosensor tip is coated with a biocompatible matrix that minimizes non-specific binding while providing a uniform and non-denaturing surface for biomolecules. BLI’s ability to characterize interactions directly in complex matrices and
non-purified samples is a key advantage. This is possible due to the robustness of the biosensor architecture and that the technology does not use complex fluidic pathways to introduce the sample to the immobilized ligand. The biosensor moves to a 96- or 384-well plate and is ‘dipped’ into the sample. This provides a robust, flexible, and a simple way to introduce an analyte to the sensor surface to monitor binding.

**Octet® BLI systems let you:**

- Acquire real-time binding kinetics data to measure the rate of association ($k_a$), the rate of dissociation ($k_d$) and affinity constants ($K_D$).
- Generate 1 to 96 data curves simultaneously with fully automated assays.
- Rapidly identify optimal conditions using up to 96 channels to assay multiple conditions and reaction configurations in a single run.
- Characterize molecular interactions in crude or unpurified matrices.
- Easily develop quantitation assays that generate data in real time.
- Detect binding of a wide range of analytes from small molecules to live cells.
- Recover precious or low-availability samples for use in another BLI experiment or other lab analyses as binding reagents are not added directly to the sample and materials are consumed minimally.

**Surface plasmon resonance**

Surface plasmon resonance is an optoelectronic method for detecting changes in refractive index at the surface of a biosensor as a result of mass changes on the biosensor surface. The biosensor is comprised of a glass substrate and a thin gold coating. Light passes through the substrate and is reflected off the gold coating. At a certain angle of incidence, a portion of the light energy couples through the gold coating and creates a surface plasmon wave at the sample and gold surface interface. The angle of incident light required to sustain the surface plasmon wave is sensitive to changes in refractive index at the surface. The change in refractive index is directly proportional to mass change that occurs as a result of molecular binding and or dissociation events. In a typical SPR assay, upon immobilization of the ligand, the analyte is supplied to the biosensor surface using a flow-based fluidics system.

Typical binding assays, whether equilibrium-based or a kinetics method, require titration of multiple analyte concentrations to accurately establish kinetics and affinity constants. While traditional SPR-based binding platforms utilize fixed concentration injections (FCI) for sample delivery, ForteBio’s Pioneer platforms offer next-generation SPR through OneStep® and NeXtStep™ injections in addition to FCI capabilities. OneStep creates an analyte gradient based on the Taylor dispersion principle in an injection line prior to entering the ligand sensing surface. The generated analyte concentration gradient consists of a full range of low to high analyte concentrations presented to the sensor surface, circumventing the need for multiple analyte titrations. With OneStep technology, a single analyte concentration gradient injection is sufficient for characterizing kinetics and affinity constants, significantly improving experimental workflows. NeXtStep injections provide a simplified way to perform competition analysis where two back to back gradient injections are performed. The first injection provides the control compound (a known binder) and the second gradient injection assesses binding of a second analyte to the control-ligand bound complex.

**SPR assays on Pioneer systems enable:**

- Acquisition of real-time binding kinetics data to measure the rate of association ($k_a$), the rate of dissociation ($k_d$) and affinity constants ($K_D$).
- High sensitivity and minimal background noise to detect small molecule and fragment binding.
- Kinetics measurements over a wide range of on- and off-rates.
- Accurate kinetics and affinity values using one analyte concentration using OneStep injections, eliminating the need to prepare a full analyte dilution series.
- High-throughput binding screens with up to 72 hours of unattended operation.

Read more about Next Generation SPR-based interaction analysis.
**BIOSENSOR ASSAYS VS. ELISA**

Kinetics characterization using biosensor-based BLI or SPR can readily replace label-dependent methods such as ELISA or fluorometric methods. Real-time biosensor assays allow one to gauge progression of the assay and also dissect binding affinities and mechanism of action with information from association ($k_a$) and dissociation ($k_d$) kinetics which ELISA does not provide.

**Benefits of biosensor assays over ELISA:**

- Biosensor assays provide binding kinetics ($k_a$) and ($k_d$) and affinities ($K_D$) compared to the end point-only affinities with ELISA.
- Kinetics assays can be completed in minutes, and real-time data allows selection of optimized association, dissociation and ligand immobilization step times as appropriate for each experiment. This avoids unnecessarily long or overnight incubation times, and enables analysis of biomolecules that are less stable under assay conditions.
- Low-affinity analytes that can often be washed away in ELISA workflows can be accurately characterized with biosensor assays.

**Benefits of ELISA over biosensor assays:**

- Low upfront costs to developing a conventional ELISA assay. A simple ELISA can be performed using conventional detection platforms such as a plate reader.
- Commonly needed ELISA reagents such as antibodies needed for capture and detection are readily available.
- Well established legacy method for quantitation and affinity determination.

**CALORIMETRY**

Calorimetry is commonly used to characterize molecular binding and structural stability of biomolecules. Isothermal titration calorimetry (ITC), allows the measurement of binding affinity ($K_D$), stoichiometry ($n$), change in enthalpy ($\Delta H$) and change in entropy ($\Delta S$) between molecules in solution by directly measuring the heat exchange that occurs during a chemical or biological reaction, such as a binding event. Differential scanning calorimetry (DSC) measures heat capacity changes ($\Delta C_p$) associated with temperature-induced unfolding of biomolecules. The resulting temperature-dependent thermogram represents a qualitative and quantitative signature of the protein unfolding. Combining both ITC and DSC data provides a holistic picture of the energetics that can help delineate varying degrees of contributions from thermodynamic properties such as entropy and enthalpy towards the reaction transformation. These parameters can also help uncover information on the binding environment such as conformational changes, solvent interactions and protonation states.

**Benefits of biosensor assays compared to ITC:**

- Affinity ranges from mM to pM can be directly measured compared to a more limited sub-mM to nM range in ITC.
- Real-time association and dissociation kinetics parameters ($k_a$, $k_d$) can be measured.
- Biomolecules from unpurified matrices can be captured to perform binding assays, allowing assays to be run in crude samples.
- Biosensor experiments require 10- to 500-fold less sample amounts for analysis compared to ITC.
- While modern ITC instruments enable users to perform up to ~20 titrations per day, biosensor assays can produce hundreds of kinetics profiles depending on the interaction system and platform, allowing them to be widely used across the drug discovery pipeline for assay development, screening, validation and characterization.
- Quantitation assays can be performed with biosensor assays.

**Benefits of ITC assay over biosensor assays:**

- Additional thermodynamic parameters for binding such as $\Delta G$, $\Delta H$, and $\Delta S$ and $\Delta C_p$ and $T_m$ (DSC) can be measured.
- Immobilization artifacts aren’t an issue as immobilization isn’t required - both binding partners interact in solution.
- No mass transfer concerns.
MICROSCALE THERMOPHORESIS

Microscale thermophoresis (MST) is another biophysical method that can be used to measure binding affinities between biomolecules. MST detects variations in fluorescent signals that result from a laser-induced temperature change. The temperature change is often generated by an infrared laser where the reaction takes place in thin capillaries. Like ITC, binding in MST is monitored in free solution, although a detectable fluorescence signal is required either by intrinsic fluorescence of the molecule (for example, tryptophan fluorescence in proteins) or generated via a fluorescent label. Intrinsic fluorescence is the preferred option as detection of the binding interaction using native conformations avoids undesirable artifacts that can arise from chemical labeling.

The MST fluorescent signal is primarily composed of two main components: temperature-related intensity change (TRIC) and thermophoresis. TRIC describes how the fluorescence intensity of a fluorophore responds to a change in local temperature. Thermophoresis describes the movement of molecules along a temperature gradient. The temperature-directed movement of molecules causes a change in the local concentrations, and the migration of a specific molecule depends on the size, charge, hydrodynamic radius and conformation. Formation of a binding complex can change the size of the assembly or cause a change in the solvation sphere, altering the thermophoretic migration pattern that can be quantitated by following the fluorescence signal. Typically, MST experiments are performed by monitoring the fluorescence of a target molecule as a function of concentration of a non-fluorescent ligand.

Key benefits of biosensor assays compared to MST:

- Kinetics parameters are obtained compared to just end-point measurements with MST.
- Labeling isn’t required, which avoids both the experimental artifacts and non-specific binding that can result from the hydrophobic fluorescent labeling molecules MST requires.
- MST is highly sensitive to any change in the binding environment including ionic strength, pH, detergents, and buffer components and require more assay optimization depending on sample conditions.
- Quantitation assays can be performed with biosensor assays.

Benefits of MST assays over biosensor assays:

- MST has a low sample volume requirement of 3-10 µL compared to a minimum 40 µL for biosensor assays.
- No immobilization is required as binding is performed in solution.

Choosing a biosensor platform

ForteBio offers two major biosensor analysis platforms, BLI-based Octet systems and SPR-based Pioneer systems, for biomolecular interaction analysis. Both offer comprehensive characterization for kinetic and affinity attributes. Biophysical laboratories can take advantage of differences in throughput, detection technologies and sample delivery between the two to establish a wide range of analysis capabilities. For example, the Octet system’s throughput and assay flexibility combined with the Pioneer system’s sensitivity to detect molecules as small as 70 Da provides a full spectrum of even the most challenging biomolecular interaction applications.

APPLICATION CONSIDERATIONS

Key attributes such as throughput, sample capacity, unattended runtimes, affinity ranges, sensitivity, type of sample matrix and ease of use are important considerations when choosing a biosensor platform to suit your application and workflow needs.
Both the Octet BLI and Pioneer SPR platforms are capable of characterizing a wide range of biomolecular interactions (Table 1). Pioneer SPR systems are ideal for small molecule and fragment screening workflows (> 70 Da) due to their exquisite sensitivity and next-generation injection capabilities. Octet BLI platforms can be used for analytes > 150 Da and applications involving larger molecular weight analytes such as viruses, nanoparticles, liposomes, and cells. Because Octet systems don’t use microfluidics, they are widely recognized as an essential tool for routine analysis of samples in crude lysates or complex matrices. Crude sample compatibility is useful especially in epitope binning, quantitation, off-rate ranking or large screening assays where sample purification is neither required or feasible for efficient workflows. Some of the key attributes of Octet and Pioneer systems are shown below for comparison.

![Image of different Octet and Pioneer SPR systems](https://www.moleculardevices.com/biologics)

Visit the website to view publications on additional applications on the Octet and Pioneer systems

<table>
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<th>Applications</th>
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<th>Pioneer FE System</th>
<th>Octet K2 System</th>
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<td>Large molecules - e.g. virus, VPL, nanoparticles, cells</td>
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<td>Screening applications - e.g. epitope binning, off-rate ranking</td>
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**Table 1**: Featured applications capabilities for Octet BLI and Pioneer SPR platforms. Recommendations for each platform is based on assay capabilities, workflow efficiency ( ), and throughput ( ) for each application. A combination of assay capability and required throughput levels should be considered when choosing a label-free instrument.
Key Octet system attributes

Throughput – Octet systems are available with anywhere from 2 to 96 channels, which directly correlates to throughput. A 2-channel system can generate data on 2 samples at the same time, and 96-channel systems generate data on up to 96 samples simultaneously. Octet K2, RED96e/QK®, RED384 and HTX systems come with 2, 8, 16 and 96 channels respectively. Channel configurations for Octet RED384 (16-channel) and HTX (96-channel) systems are shown in Figure 2. The Octet HTX system can adapt to a user selectable 8, 16, 32, 48 or 96-channel read-head to tailor assay design. It can also complete a 96-well plate quantitation in 2 minutes, a full plate of off-rate ranking assay in minutes, and a 32 x 32 epitope binning screen in less than 8 hours for example. The 8-channel Octet RED96e and QK® systems are widely used for kinetic characterizations, quantitation assays and medium-throughput screening campaigns. The Octet K2 system is ideal for labs with lower throughput needs that also require high-performance biomolecular characterization capabilities at an economical price point. While all Octet platforms are fully automated, unattended operation can be further extended with Octet RED384 and HTX systems by integrating tailored robotic configurations for advanced assay workflows.

Learn more about Octet systems.

Sample capacity – Samples and reagents are housed in 96- and 384-well plates. Octet K2, RED96e, and QK® systems hold one 96-well plate and Octet RED384 and HTX systems have two plate positions where either 96- and/or 384-well plates can be used. A 200 µL minimum sample volume is required for standard 96- well plates and this can be reduced to 40 µL when using 384 tilted-well plates. The increased sample capacity and flexibility with plate types for assays on Octet RED384 and HTX systems allow lower sample volumes to be used when required, and enable analysis on more samples. For complex assay workflows that require multiple reagents, this increased sample capacity can also be beneficial in that the number of user interventions during the assay can be minimized.

Broad range of affinities and kinetics – Octet systems capture a wide range of kinetic association ($10^{-1} – 10^{7} M^{-1}s^{-1}$) and dissociation ($10^{-6} - 10^{-1} s^{-1}$) rates and measure affinities ($K_D$) from 1 mM – 10 pM, representing biologically relevant interactions.

Small and large molecule analysis – Octet platforms analyze a wide molecular weight range, from small molecules (> 150 Da) to large nanoparticles, viruses and cells providing the capability to characterize a multitude of interaction systems using a single platform.

Learn more about robotic automation of Octet assays for high-throughput ligand screening.

Octet RED384

Octet HTX

Figure 2: Biosensor channel configurations in Octet systems.
Sample delivery – Sample introduction with Octet systems is accomplished by dipping biosensors directly into the samples, no fluidics are used. This provides greater flexibility in sample preparation and use of different sample types with none of the concerns of clogged fluidics associated with other methodologies.

Walk-in ready – No prior instrument preparation such as cleaning, priming or equilibration is required to run assays on Octet systems. Simply design the experiment, load samples and run.

Key Pioneer system attributes

Throughput – Pioneer and Pioneer FE systems have three flow cells, two of which are typically used for detecting interactions, with the third being used as a reference. This allows kinetic data from two ligand targets to be generated simultaneously.

Simplified workflows – Unlike traditional kinetics and affinity analysis workflows, Pioneer systems don’t require a full dilution series with multiple analyte concentrations to characterize a binding interaction. Unique to these systems, OneStep Gradient Injections provide a continuous analyte concentration gradient of 3–4 orders of magnitude in a single injection. This reduces the number of samples that have to be analyzed (Figure 3).

Learn more about OneStep Injection technology.

Sample capacity – Samples and reagents are introduced to the biosensor surface using a tightly-controlled automated fluidic system. Pioneer systems can hold up to two 96- and/or 384-well plates with two reagent racks providing the capacity to analyze up to 768 fragments in a single unattended assay.

Broad range of affinities and kinetics – Pioneer systems can capture a wide range of kinetic association ($10^{-2} - 10^9 \text{ M}^{-1} \text{s}^{-1}$) and dissociation ($10^{-6} - 2.5 \text{ s}^{-1}$) rates and measure affinities ($K_D$) from 1 mM – 1 pM.

Small and large molecule analysis – Pioneer systems provide the sensitivity and baseline stability needed to detect binding interactions for a wide range of analytes, from biologics to small molecule fragments as low as 70 Da.

TYPICAL EXPERIMENTAL WORKFLOW

Despite differences in detection and sample delivery, experimental workflows for measuring kinetics and affinity constants between BLI- and SPR-based systems is similar (Figure 4). The main assay steps include:

1 Ligand loading (immobilization) to the biosensor:
   Immobilization of one binding partner using a suitable biosensor chemistry. For example, a His-tagged protein can be immobilized using a Ni-NTA biosensor, or biotin-labeled proteins or nucleic acids can be immobilized to a streptavidin-coated biosensor. It’s critical to choose a suitable immobilization chemistry to present the ligand favorably towards the analyte molecule.

   See our BLI and SPR biosensor chemistry selection guides for a full list of available biosensors.

2 Post-immobilization baseline:
   After immobilization, the ligand is introduced to binding/assay buffer to equilibrate and wash away loosely bound ligand molecules.

3 Analyte association:
   Analyte is introduced for binding analysis. Association kinetics are measured in this step. While the analyte concentration is required for calculating association rates, in the absence of this information, the presence of a binding (Yes/No) interaction can be tested. Octet systems can also utilize the association phase for quantitation purposes in lieu of ELISA.

4 Analyte dissociation from the ligand:
   The bound complex is dissociated in assay buffer. Unlike the association process, the dissociation is not dependent on reagent concentrations. This enables accurate calcu-
Figure 4: Typical assay steps in a kinetic binding experiment to measure binding kinetics (association and dissociation) and affinity constants.

5 Regeneration: After data collection, the regeneration process strips the bound analyte from the immobilized ligand which prepares the surface for another round of analyte binding. For ligands immobilized by affinity capture methods, for example with Ni-NTA or Protein A biosensors, regeneration usually removes the ligand molecule as well. In these scenarios, re-immobilization of the ligand is required prior to analyte binding.

The total experimental time for generating a kinetic curve depends on the time of each individual step, which in turn depends largely on the kinetic properties of the interaction itself. An interaction system with a rapid on-rate can reach equilibrium within a few minutes, a slow dissociating complex can require a much longer time at the dissociation phase. For a typical interaction, an estimated time for a kinetic trace would be about 5–30 min of the total experimental time. It’s also well established that to accurately characterize interactions, assaying binding at multiple analyte concentrations is required, preferably below and above $K_D$ concentrations (Figure 5). Kinetic curves are then fitted as a group (global fitting) to calculate for kinetics parameters. While traditional biosensor assays would require generating multiple kinetic curves representing several analyte concentrations, the Pioneer SPR platform only requires one kinetic curve to characterize kinetics and affinity values.

Learn how to develop a kinetic binding assay on the Octet platform.

Learn how to develop an OneStep SPR assay using Pioneer SPR systems.

Figure 5: Analyte titration series to calculate kinetics and affinity constants. A six point analyte titration (blue) and 1:1 binding model fits (yellow) are shown. Data were fitted using Octet Data Analysis software by global fitting unifying all kinetic traces to fit for association ($k_a$), dissociation ($k_d$) kinetics and affinity ($K_D$).
Featured applications

ANALYSIS ON FC-GAMMA RECEPTOR INTERACTIONS IN OCTET PLATFORMS

Fc gamma receptors (FcγRs) are membrane glycoproteins found on certain types of cell surfaces that contribute to immune system functions. With high binding specificities to Fc regions of immunoglobulin G (IgGs), Fc-receptors bind to IgGs displayed on pathogenic cells to trigger antibody mediated cell-dependent cytotoxicity (ADCC) or phagocytosis. Fc-receptor interactions bind IgGs with a broad range of affinities based on the type of receptor involved. FcγRI (CD64) affinities to IgGs are characterized at 0.1 nM–10 nM, while FcγRII (CD32) and FcγRIII (CD16) interactions are measured at 0.1–10 µM affinities. In addition, binding properties can be affected by genetic polymorphisms of the receptors as well as glycosylation patterns in the Fc region on the antibody.

Octet BLI platforms are utilized for characterizing Fc-gamma receptors due to their:

- Ability to rapidly measure binding interactions across a wide range of affinities (1 mM–10 pM).
- Detect variabilities in affinities due to variations such as glycosylation patterns.
- Availability of a wide range of biosensor chemistries for ligand immobilization.
- High-throughput, and lower consumable costs.

An example workflow for performing an FcRn-hIgG kinetic assay is shown in Figure 6.
Figure 6: Workflow of a FcRn-hIgG kinetic assay using anti-human FAB2G biosensors. (A) hIgG is immobilized and FcγRIIIa is introduced as the analyte. Other commonly used biosensor chemistries are Ni-NTA and Streptavidin. (B) Kinetic analysis using a 1:1 interaction model to determine kinetics and affinity. (C) Steady state analysis using data represented in (B). Steady state (end-point) analysis can also be useful especially in low affinity interactions or for systems that produces very fast on-rates.

<table>
<thead>
<tr>
<th>% Deglycosylation</th>
<th>$K_D$ (M)</th>
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<td>0</td>
<td>8.68E-07</td>
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<tr>
<td>10</td>
<td>9.89E-07</td>
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<tr>
<td>25</td>
<td>1.13E-06</td>
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<tr>
<td>50</td>
<td>1.52E-06</td>
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</table>

Figure 7: Binding of FcγRIIIa is affected by glycosylation and can be measured by an Octet binding assay. Monoclonal hIgG with various levels of glycosylation profiles binding to FcγRIIIa. Interaction becomes weaker with reduced glycosylation levels.

Learn more about analyzing Fc-gamma receptor-IgG interactions.

Learn more about analyzing FcRn-antibody interactions.
**QUANTITATION ON OCTET BLI PLATFORMS**

Octet platforms offer an excellent solution to replace assays that are currently performed as ELISA. Similar to ELISA, Octet assays are carried out on an immobilized surface (solid support in ELISA), and the analyte is bound from solution. However, Octet assays are fully automated, require much less user intervention, and provide a simplified workflow. In addition, they provide researchers with the flexibility to choose the most efficient assay format based on assay needs such as sensitivity, dynamic range, and workflow (Figure 8).

Advantages of Octet quantitation assays over ELISA:

- Multiple assay formats – depending on the required sensitivity of the assay, several assay formats can be adapted: a single step, sandwich or the 3-step assay which provides the highest sensitivity.
- Quantitation assays are easily developed for a variety of analytes such as IgG, recombinant proteins, vaccines and viruses, and more.
- Detection of low affinity analytes that can often be washed away during ELISA procedures due to their faster off rates.
- Real-time data generation to assess experimental progression.
- Faster time to results which is advantageous in detecting less stable analytes.
- Samples and reagents can be fully recovered and reused.
- Lower assay costs as biosensors can be regenerated.

Learn how to convert your ELISA assays to Octet assays.

**Selecting an assay format**

The choice of an assay format is dependent on the concentration range of analyte to be quantified (Figure 8). Direct binding quantitation assays (1-step) typically can measure concentrations from low mg/mL up to low ng/mL depending on the analyte. Larger analytes provide higher sensitivity compared to lower molecular weight analytes.

1-step assays are simpler, faster and eliminate the need for secondary reagents and steps needed for detection.

<table>
<thead>
<tr>
<th>Assay features</th>
<th>1-step</th>
<th>2-step</th>
<th>3-step</th>
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<tr>
<td>Pictorial representation</td>
<td><img src="image1" alt="Pictorial representation of 1-step assay" /></td>
<td><img src="image2" alt="Pictorial representation of 2-step assay" /></td>
<td><img src="image3" alt="Pictorial representation of 3-step assay" /></td>
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<tr>
<td>Typical assay time</td>
<td>• 30 min (Octet QK®, RED96e) • 15 min (Octet RED384) • 2 min (Octet HTX)</td>
<td>• 1 hr 30 min (Octet QK®, RED96e) • 1 hr 15 min (Octet RED384) • 1 hr (Octet HTX)</td>
<td>• 2 hr (Octet QK®, RED96e) • 1 hr 30 min (Octet RED384) • 1 hr (Octet HTX)</td>
</tr>
<tr>
<td>Typical concentration range</td>
<td>Low mg/mL to low ng/mL</td>
<td>Low ng/mL to low pg/mL</td>
<td>Low ng/mL to low pg/mL</td>
</tr>
<tr>
<td>Advantages</td>
<td>• Single incubation step — fast, easy, reduces reagent expenses • Low affinity analytes detected — even those missed by ELISA • NO labeled reagents • Kinetic parameters can be measured • Allows regeneration and re-use of biosensor in most cases</td>
<td>• Two incubation steps — still fast, easy, reduces reagent expenses in comparison to ELISA • Higher sensitivity of detection, down to low pg/mL, depending on assay • NO labeled reagents • Automated and no-wash assay minimizes handling</td>
<td>• Similar to most ELISA assays in format — but faster and easier • Excellent sensitivity — down to low pg/mL, depending on assay • Automated and no-wash assay minimizes handling</td>
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**Figure 8:** Octet quantitation assay formats and features.

[www.moleculardevices.com/biologics](www.moleculardevices.com/biologics)
In addition, they enable researchers to reuse biosensors after regeneration in most cases. Multi-step assays (2- and 3-step) are used when enhanced sensitivities are required up to low pg/mL concentrations (Figures 9 and 10). Since Octet detection relies on the optical thickness generated at the biosensor, increasing the binding layer by 2 or 3 step analytes increases the binding signal and sensitivity. The 2-step format typically includes the capture of the analyte by the biosensor followed by sandwiching the analyte by a second antibody. For higher signal amplification, an enzyme-linked sandwich assay (3-step assays) captures analyte bound by two separately-labeled capture molecules to the biosensor in the first step, binds an HRP-conjugated antibody to the complex in the second step, and precipitates a substrate directly onto the biosensor surface further increasing the mass added in the third step. Read about how you can build you own custom quantitation assays.

**Figure 9:** Sandwich assay to quantitate Protein A. Protein A is widely used in purification columns to capture and purify IgGs. However, during the purification process, Protein A can leach off the chromatography matrix and co-elute with IgGs. A sandwich-based (2-step) assay can be used to quantitate protein A co-eluted with IgG. (A) BLI sensorgrams for Protein A standards are generated using protein A concentrations from 25 ng/mL to 0.1 ng/mL (multiple replicates). (B) Binding rates from (A) are plotted for each protein A standard concentration. The standard curve points are fitted to a 4-PL model in Octet data analysis software.

**Figure 10:** Quantitation of recombinant insulin in DMEM media using a 1-step Octet quantitation assay. (A) BLI sensorgrams for insulin binding to immobilized biotin-labeled anti-insulin antibody at concentrations of 0.78–50 μg/mL. (B) Standard curves generated by plotting initial binding rates from (A) against standard concentrations. Data were fitted using a 5-PL dose-response curve fitting equation in Octet Data Analysis software replicates). (B) Binding rates from (A) are plotted for each protein A standard concentration. The standard curve points are fitted to a 4-PL model in Octet data analysis software.
Epitope binning assays also are performed to identify mAbs that bind similar epitopes to a previously characterized mAb for generations of biosimilars or biobetters. These assays can also be useful in reagent development programs to identify antibody pairs that are best suitable for development of ELISA assays.

Octet systems provide several advantages for these assays:

- High-throughput and assay speed.
- Versatility - choose from three assay formats (Figure 11).
- Availability of a variety of biosensor chemistries suitable for analysis.
- Ability to characterize antibodies from hybridoma supernatants, phage lysates or purified samples.

Learn more about epitope binning assays on Octet platforms.
**FRAGMENT SCREENING WITH ONESTEP SPR**

Fragment based drug discovery (FBDD) approaches are aimed at identifying chemical scaffolds that interact with a target molecule and alter or inhibit activity. Typical fragment libraries consist of compounds with 6–15 heavy atoms presenting low complexity scaffolds resulting in low affinity binding interactions that are in the range of 10 µM to 10 mM. Sensitive detection technologies are required to successfully capture fragment binding hits. The Pioneer FE SPR platform is an ideal solution to establish fragment binding screens due to high sensitivity, throughput and the availability of OneStep and NeXtStep injections that significantly simplifies and reduce screening workflow timelines.

In Pioneer FE system fragment screening projects, using a single concentration copy of the fragment library researchers are able to:

- Determine kinetics ($k_\text{on}, k_\text{off}$) and affinity ($K_d$) constants from the primary fragment screen with OneStep® Injections (Figure 14).

![Learn more about fragment screening on the Pioneer FE system](image)

**Figure 13**: Data analysis. Two-dimensional matrix for a panel of 64 x 64 mAbs profiled in an in-tandem binding assay (4096 total pairing reactions). The saturating antibodies (Ab1) and competing antibodies are listed in rows and columns respectively. Red indicates a blocking interaction (competing) pair and green indicates a binding pair. The matrix diagonal represents self-self interactions for each mAb that is highlighted in orange. Related antibodies were clustered into bins using a Pearson clustering analysis. Data was analyzed using Octet Data analysis HT version 11 software that includes a dedicated module for unifying multiple runs and analysis of large epitope binding data sets.

**Figure 14**: OneStep fragment screening using the Pioneer FE system. SPR sensorgrams from a fragment screen are shown. Blue curves represent the positive control compound binding that run intermittent-ly to assess protein activity throughout the screen. The inset depicts equilibrium response values for each fragment vs. cycle number. The hit selection feature (Pioneer FE only) in Qdat software enables identification of fragment hits and calculates kinetics and affinity values.

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• Test up to 768 fragments in 24 hrs with OneStep and eliminate secondary screening that involves dose-response titrations.
• Test up to two therapeutic targets versus a fragment library simultaneously.
• Test fragments for competition mechanism and determine kinetics ($k_a$, $k_d$) and affinity ($K_D$) in the presence of a competitor molecule using NeXtStep™ Injections.
• Accurately select hits from primary screen data using a novel normalization and selection algorithm developed in collaboration with industry leaders.
• Gain significant time and reagent savings compared to conventional SPR-based fragment screening.

CHARACTERIZING MECHANISM OF BINDING USING NEXTSTEP COMPETITION ANALYSIS

Competition binding assays can be used to identify site-selective binding or gain more information on the binding mechanism of the identified hits. The Pioneer FE system utilizes NeXtStep injections to generate full kinetics and affinities in the presence of a competitor. NeXtStep consists of two back to back gradient injections, in which the first injection results in the formation of the binding complex with a known compound (positive control) at saturating conditions. Once all binding sites are occupied, the second gradient is established with the fragment. If the fragment requires the identical binding site as the positive control, no binding or reduced response can be expected (competitive inhibitors, Figure 15B), while an unperturbed binding response would indicate a non-competitive binding mechanism (Figure 15A).

CHARACTERIZING IRREVERSIBLE INHIBITORS USING PIONEER SPR ASSAYS

The majority of inhibitors characterized on biosensor platforms are under equilibrium binding conditions where the inhibitor-ligand complex formation is rapid and reversible. However, ~30% of the approved therapeutic enzyme inhibitors in the market function through covalent modification of the target. While toxicity, insufficient efficacy, clearance, and metabolite reactivity are commonly cited pitfalls for covalent inhibitors, approved therapeutics in this space show high selectivity and potency towards the binding target that drives overall therapeutic efficacy.

Analysis of covalent inhibitors in SPR can be challenging due to inactivation of the immobilized target by covalent inhibitor binding and the inability to reuse the surface for subsequent inhibitor analyses. Affinity capture methods such as His-tag capture immobilizations enable reversible target immobilizations that can be suitable for reversible inhibitor analysis, but these methods are often associated with drifting baselines, which can affect data analysis. A reversible streptavidin-biotin capture method can be adapted in Pioneer SPR systems that is suitable for irreversible inhibitor analysis (Figure 16).

In contrast to conventional interactions, covalent inhibitors operate under non-equilibrium kinetics and drive towards complete saturation of the target site if sufficient contact time is provided. Therefore, covalent inhibitors cannot be efficiently ranked using traditional IC50 measurements and instead require a consideration of the rate of inactivation of the target molecule. An analytical model describing the kinetics of inactivation is shown in (Figure 17) that is part of the Pioneer SPR data analysis suite. Here, a commitment to covalency (Cc) parameter is introduced which relates to the efficiency of the adduct (E-I) formation and its relationship to dissociation. Together, efficiency of adduct formation and the rate of protein/inhibitor association defines the biochemical potency of the inhibitor.

Learn more about characterization of irreversible inhibitors on Pioneer SPR systems.

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**Figure 15:** NeXtStep injections with the Pioneer FE system. Binding profile in the presence of a control (blue) and binding profiles in the absence of control (black). (A) Noncompetitive fragment binding shows an unchanged binding response in the presence of a control. (B) Fragments with competitive binding show decreased binding responses in NeXtStep injections in the presence of the control.
Figure 16: Amine-PEG4-desthiobiotin is coupled to an amine reaction biosensor to create a desthiobiotin sensor surface. The biotin-labeled target is then captured using streptavidin and immobilized onto the desthiobiotin sensor surface for analysis with covalent inhibitors. The target-inhibitor complex is then regenerated using an acetonitrile:NaOH mixture. The surface can then be re-immobilized with fresh target for multiple rounds of covalent inhibitor analysis.

Figure 17: Analytical model for kinetics of target inactivation by irreversible inhibitors. (A) Rate equations for irreversible inhibition, where E is enzyme (target), I is inhibitor, E•I is encounter complex and E-I is adduct or covalent complex. $k_{on}$ and $k_{off}$ are rate constants of association and dissociation respectively. $k_{inact}$ is the reaction rate constant of covalent inactivation. (B) Kinetics of inactivation is described. $K_i$ and $K_I$ describe inhibitor potency for reversible and irreversible inhibition, respectively. $k_{inact}$ is the rate of covalent reaction to form the adduct, E-I. By this model both affinity ($K_i$) for the target, as well as highly efficient chemistry ($k_{inact}$) are required to get efficient irreversible inhibition. The Cc term introduced is the “commitment to covalency” representing the probability that the encounter complex proceeds to form the adduct. The product of Cc and the association rate constant, $k_{on}$, describes the overall biochemical potency ($k_{inact}/K_I$).
Figure 18: SPR data from the Pioneer FE system showing three covalent kinase inhibitors binding with different $C_c$ values. Signal after referencing shows stable baselines, suitable for measuring these potent inhibitors and differentiating binding from covalent reaction. Potency parameters are compared between inhibitors and biochemical assay methods to show the differences in reaction efficiency.

### CHARACTERIZATION OF HIGH AFFINITY BIOLOGICS INTERACTIONS WITH ONESTEP ON PIONEER SPR SYSTEMS

Recent advances in protein engineering technologies have contributed to the development of several breakthrough biologics to treat diseases such as cancer, inflammation, and diabetics. Biologics or protein therapeutics often can provide higher target specificity and reduced side effects compared to small molecule drugs. Today’s biologics drug discovery process involves a hit-to-lead campaign where lead candidates are identified by screening thousands of protein samples using biochemical, ELISA or binding screens. Octet BLI systems are often used in these screens due to their compatibility with crude lysate testing. Lead identification is followed by target binding characterization, an essential step in lead validation, to select for highly selective biologic candidates with desired kinetics and affinities towards the target.

Pioneer Next Generation SPR platforms are ideal for high-affinity lead characterization and validation due to:

- **OneStep characterization is 3- to 6-fold faster than a traditional characterization on a 4-channel SPR system.**
- **OneStep injections use less sample (50% less analyte).**
- **No analyte dilutions are required.**

With OneStep gradient injections, binding kinetics and affinity values can be measured using a single analyte titration step which significantly improves the efficiency of characterization workflows over traditional SPR. Since most high-affinity interactions produce slow off-rates (dissociation) from the target, longer dissociation analysis times are required for accurate kinetic analysis. When traditional SPR workflows are utilized that require multiple analyte concentrations per target the experimental times can be significantly longer compared to OneStep, where a single titration is sufficient for analysis (Figure 19).

<table>
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<th>Published inhibitors</th>
<th>$k_{on}$</th>
<th>$C_c$</th>
<th>$k_{inact}/K_i$ (M$^{-1}$s$^{-1}$)</th>
<th>$k_{inact}/K_i$ (M$^{-1}$s$^{-1}$)</th>
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<tr>
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<td>7.8E+04</td>
<td>2.1E+05</td>
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<tr>
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<td>0.6</td>
<td>5.5E+05</td>
<td>7.3E+05</td>
</tr>
<tr>
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<td>1</td>
<td>4.1E+05</td>
<td>8.8E+05</td>
</tr>
</tbody>
</table>
Figure 19: OneStep characterization of a panel of eight mAb candidates. (A) Eight mAb hit candidates binding against a target protein are assayed using the Pioneer FE SPR system. Binding data (black) are least-squares fit to a 1:1 kinetic binding model (red). Total experimental time for OneStep binding assays is ~20 hrs hours including buffers controls. The identical experiment with seven analyte titrations per target would require ~63 hrs on a traditional SPR system with fixed concentration injections. (B) Affinity plot ($k_d$ vs. $k_a$) for target binding to mAbs characterized by OneStep SPR. Dashed lines indicate common affinity relationships with different kinetic rate constants.

Learn more on OneStep characterization of high-affinity biologic interactions.

Figure 20: Pioneer systems provide excellent baseline stability for accurate kinetic characterizations of high affinity binding interactions. This is indicated in the OneStep characterization of a high-affinity binding interaction with extended dissociation times (up to ~5 hrs). Figure inset show the first 2000 sec of data including the full OneStep association (0–400 sec) followed by the first 1600 sec of the dissociation phase.
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Octet K2 System

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Octet RED384 System

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Pioneer FE System