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Octet® SPR Analysis Features

This tutorial outlines the operation of Octet® SPR Analysis software, a data analysis software package that has been tailored for Octet® SF3 data sets. The program allows rapid processing of binding response curves for kinetics and affinity analysis. The intuitive, user-friendly interface guides the user through a series of data transformations that produce data sets that are then fitted with standard kinetic or affinity models. A key feature is the ability to rapidly perform single or double referencing of the data set in advance of model fitting.

When using solvents such as Dimethylsulfoxide (DMSO) a simple built in tool enables correction of the excluded volume effect.

Kinetic and affinity models in Octet® SPR Analysis software include:

- Simple off rate model for determining the dissociation kinetic rate constant \(k_d\)
- Pseudo-first-order kinetic model (1:1 interaction) for determining the association \(k_a\) and dissociation kinetic rate constant \(k_d\)
- Two compartment model (mass transport limitation)
- Bivalent analyte model
- Two-state model
- Gradient injection specific models for diffusion and protein aggregation

In addition to kinetic models, a solid phase steady-state affinity calculator is included for use with steady-state binding data.

A versatile graphics interface permits viewing of the data in a variety of formats with customizable parameters including fonts, colors and scaling. Where desired, data may be easily exported as columns of text for plotting in user-defined data analysis programs or as publication ready image files that may be pasted into your word processing software.

Terminology

SPR instrumentation follows the course of an interaction between a molecule bound to the sensor surface (ligand) and a second free, solution-phase molecule (analyte).

Table 1-1: Key Terms

<table>
<thead>
<tr>
<th>Term</th>
<th>Description</th>
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<tr>
<td>Ligand</td>
<td>The interaction partner attached to the surface of the sensor, In screening applications, this may also be referred to as the Target.</td>
</tr>
<tr>
<td>Analyte</td>
<td>The solution-phase interaction partner that is injected over the surface-bound ligand.</td>
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</table>
Opening Octet® SPR Analysis

Octet® SPR Analysis can be opened via a number of routes:

- By clicking on the desktop shortcut
- By clicking on the Analyze button located on the View Page of the Octet® SPR Discovery software (Figure 1-1)
- By clicking the Octet® SPR Analysis shortcut located in the Octet® SPR program group

![Figure 1-1: Octet® SPR Analysis Application Screen](image)

A splash screen will appear while the Octet® SPR Analysis application opens.

The Octet® SPR Analysis application consists of a series of tabbed pages, each of which performs a transformation of the data set. The objective of Octet® SPR Analysis is to rapidly perform reference curve subtraction and kinetic or affinity analyses with minimal data manipulations. In some cases, Octet® SPR Analysis will automatically perform these transformations if the Octet® SPR Discovery protocol included the default settings for analysis.

![Figure 1-2: Tabs for Response Curves](image)

Response Curve Selection

Specific response curves from experimental data can be viewed by selecting from one of the drop-down boxes seen in Figure 1-2: Assay Filter, Analyte Filter, or Target Filter. These filters default to show all data but can quickly isolate data of interest for review.

Color coding of the data can be adjusted by changing the selection of the Curve Coloration drop-down box. The color codes are predetermined by the software and the available options are: Color by Concentration, Color by Assay, Color by Analyte, Color by Target and Black (default).
Data can also be scaled according to analyte molecular weight (MW) by clicking the **Correction Factor: MW** check box. Data files exported to Octet® SPR Analysis through the Octet® SPR Discovery control software will automatically transfer analyte MW, if available. The median MW will be identified and all responses normalized to this MW. Therefore, theoretical $R_{\text{max}}$ calculations can be made using this analyte MW. A similar normalization can be applied based on the analyte refractive index (RI) by selecting the **Correction Factor: RI** check box.

### Despiking

Data opened in Octet® SPR Analysis are automatically Despiked, meaning that an algorithm scans the data and removes/smooths a window of data around perceived spikes. This setting can be turned off by clicking the **Configure** menu button and unchecking **Despike**. The Despike settings can be adjusted by clicking **Despike options**. The Number Peaks Tested defines the maximum number of spikes that can be identified and corrected in a data set. The Peak Window Size adjusts how much data is reviewed to identify and correct a spike. Larger Peak Window Sizes will extract larger spikes and lower Window Sizes will extract a more refined spike. The Spike Threshold establishes the minimum jump in response that should be considered by the algorithm as a potential spike. Figure 1-3 shows data before (left panel) and after (right panel) Despike.

![Figure 1-3: Spike (left) and Despike (right) Response Curves](image-url)
Chapter 2:
Data Tab

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Data Page Description

The Data tab is the initial user interface for review of absolute, unreferenced SPR data.

If not accessed through Octet® SPR Discovery software, click on the Data tab, click the Open button on the toolbar and load the desired data (Figure 2-1). Octet® SF3 data files appear as .spr files. File types used by Octet® SPR Analysis are .qdtx (raw SPR data export) and .qdt (analyzed data files). A simulated data set for a monovalent interaction is used to illustrate model fitting. In this experiment, serial tripling dilutions of protein were analyzed by injecting them sequentially over all three sensing channels where flow channel (FC) 1 was coated with target.

![Image of Data tab with file selection dialog]

**Figure 2-1: Opening the Data Tab**

The data set is shown in the plot window in Figure 2-2.
The overlaid binding response curves shown in the plot are response curves for all data channels. In general, the three channels are represented; two channels immobilized with a specific ligand or capture molecule for kinetics and affinity assessment and an appropriate reference channel. High-quality data requires processing the data with use of this reference surface. Many of the artifacts associated with SPR binding data can be removed by proper experimental design and an appropriate reference surface is critical. In this example the simulated data are identical between channels and a true reference channel was not present.

**Entering and Correcting Analyte Concentrations and Names**

**Correcting analyte concentrations**

A sample table appears on the left-hand side of the plot window with a series of analyte and target information (Figure 2-3). Each row represents three curves in the plot window, namely the binding response curves for the interaction of the analyte with each of Octet® SF3’s three flow channels. The concentrations shown in the sample table for each cycle are the analyte concentrations entered for each cycle when recording the data using the Octet® SPR Discovery software.

If the concentration was entered incorrectly in the Octet® SPR Discovery software when preparing the experiment, the correct concentration can be typed into the appropriate cell in the table. By checking the **Show Individual Curves in Table** option, the information for target versus cycle can be modified. In most cases this will be constant but for certain capture assays the Target may change per cycle.
The analyte concentrations are automatically entered terms of Molarity (M). Alternative abbreviations include:

- m (10^{-3} M)
- u (10^{-6} M)
- n (10^{-9} M)
- p (10^{-12} M)

For example, where correcting the analyte concentration these can be typed as either “1u” or 1e-6 for a 1 µM sample. This method of entering the concentrations is equivalent to the earlier table but is sometimes more convenient, especially when changing a small number of samples.

Note that the table recognizes either designation and can work with a mixed table containing both concentration designations.

Correcting analyte names

If an Analyte name was incorrectly entered, it can be changed using the Reassign feature found by right-clicking on an Analyte name. The analyte name can be chosen from the list of pre-existing analyte IDs or if not present in the list can be entered manually in the New text box (Figure 2-4).
Information from the Analyte table can be copied to the clipboard by right-clicking in the table and clicking **Copy Table**. The table information can then be pasted and/or modified for more intensive sorting or name change transformations. Then the table can be re-pasted by right-clicking the Octet® SPR Analysis Analyte table and clicking **Paste Table** (Figure 2-5).
Plot Zoom

The plot window scales according to the upper and lower limits of the data set. However, the response for the actual binding phase of these curves is often much lower than the range of the y-axis, for example when a high-refractive index regeneration is used. It is therefore often helpful to zoom in on the binding region using the zoom function. Start by drawing a box around the region of interest by clicking and dragging the left mouse button. Next right-click and choose the **Zoom In** option from the pop-up menu (Figure 2-6).

![Figure 2-6: Zoom Function](image)

A magnified view of the selected region appears in the plot window (Figure 2-7).

![Figure 2-7: Zoomed Region](image)
Eliminating Poor Data Sets

A response curve may be removed from the analysis, for example, if it can be clearly identified as an outlier, or is of poor quality.

A response curve is removed by one of three methods:

- If the cycle number of the injection to be hidden is known, it can be removed by clicking the Hide check box in the Data tab sample table. A red highlighted curve on this page indicates a hidden curve which will not appear in subsequent graphs.

- Clicking the Hide icon found at the tool bar in the top left side. This dialog enables sorting of response curves in a variety of ways and show/hide are options for any of these data views.

- Double-clicking the curve to be hidden and then right-clicking in the graph to choose the Exclude line option. The associated cell in the sample table will be grayed out and the associated curves for all three channels will appear red in the Data page plot and be removed from all subsequent tab pages.

![Figure 2-8: Eliminating Data Sets](image)
Chapter 3:

Prepare Tab

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Description of the Prepare Tab

The Prepare tab contains all the components for data preparation including double reference subtraction, aligning and calibrating.

The data analysis steps progress from left to right: Zero, Crop, Align, Reference, RI Calibration, and Blanks. The Prepare tab shows three views of the data which change when a particular Prepare step is clicked:

- The before graph on the left shows the view of the data before a particular Prepare step has been applied.
- The after graph in the middle shows the data after the particular Prepare step has been applied.
- The final graph on the right shows the data as it will be after all Prepare steps have been applied. Clicking in one of the Prepare step boxes will change the focus of the left and middle graphs to the changes associated with that step.

![Prepare Tab](image)

**Figure 3-1: Prepare Tab**

The following sections will expand on the features within each Prepare step.

**Zero Step**

The Zero step allows the set of response curves to be y-normalized so that a zero baseline is set for all curves in the set. The left (green) and right (red) time limit lines are positioned near to the analyte injection phase of the curves. Usually a 5-10 second window, as delimited by these vertical time lines and located immediately before the injection phase, is appropriate. The left plot (Figure 3-2) shows the positions of the selected vertical time lines. The selected times are displayed in the Average from and to boxes in the zero tab. Note that the injection start point defaults to a time of zero seconds. When the Zero check box is checked, the transformed plot appears on the middle plot. Note that absolute responses (RU) vary in the before graph, this is due to the presence of an immobilized protein on only one flow channel and a significant reason for the Zero adjustment of the data.
Crop Step

The Crop step allows the elimination of unwanted regions of the response curve before reference curve subtraction and model fitting. It is recommended to include at least 30 seconds of baseline data before the injection and dissociation phase while excluding sensor wash steps or the regeneration phase (if applicable). Place the green time line on the baseline at least 30 seconds before the injection phase. Place the red time line just before the regeneration phase. Check the Crop check box. The selected data set between these limits is then shown in the right hand plot.

Figure 3-2: Zero Step
window. The data set now consists of only the regions required for model fitting, i.e. $k_a$ (association) and $k_d$ (dissociation).

Figure 3-3: Crop Step
An incorrectly cropped data set is shown below (Figure 3-4) where a section of the association phase has been cropped out. This will interfere with model fitting and must be corrected by adjusting the start time (green line) before the start of injection. This is a common error when analyzing OneStep® data where the beginning of the injection is important for the concentration vs. time profile.

Figure 3-4: Uncorrected Cropped Data Set

**Align Step**

Octet® SF3’s computer-controlled fluidic design ensures that the timing of injections from one cycle to the next is highly reproducible. In addition, injections are time-stamped, allowing automatic alignment of injection start times when exporting to Octet® SPR Analysis. In cases where large refractive index offsets are present in the injected samples, it may be necessary to correct for channel time-of-arrival differences using the channel-to-channel Align step. The Octet® SPR Discovery software is able to automatically estimate this correction for certain assay types. Each channel may be adjusted independently forward or backward in time (preferably leaving one channel at 0). See Figure 3-5 for an example.

It is recommended to observe the bulk refractive index response of a non-binding molecule such as DMSO when adjusting channel-to-channel alignment in order to achieve proper referencing.

Figure 3-5: Setting Align Step Parameters
It is possible to have each injection of a manual injection curve broken into its own cycle for purposes of analysis. Begin by loading the data file into the View tab of the Octet® SPR Discovery Software. After clicking **Analyze** in the Octet® SPR Discovery Software, check the box **Break into one cycle per injection** (see Figure 3-6). This feature is built into the Octet® SPR Discovery software if all injections were performed in the same response graph. Refer to the Octet® SPR Discovery User Guide for further instructions.

![Export Option: Break Into One Cycle Per Injection](Figure 3-6: Export Option: Break Into One Cycle Per Injection)

**Reference Step**

Use the Reference step to subtract the interaction of analyte with the reference sensing surface from the response. In the Reference Step, channel 2 contains the reference curves. After using the Reference Step, the corrected data set is displayed in the right hand plot. This transformation has eliminated the small response steps due to bulk refractive index offsets at the beginning and end of each injection. Any non-specific binding or baseline drift due to other
interferences are also subtracted, making this a powerful means of isolating the responses due to analyte binding in the presence of artifacts.

![Figure 3-7: Reference Step](image)

**RI Calibration Step**

The process outlined below can be simplified by using the Assay Micro-calibration feature. More information can be found on this technique in the Octet® SPR Discovery User Guide.

Dimethylsulfoxide (DMSO) is a common solvent included in SPR assays with small molecule analytes. However, DMSO possesses a high refractive index and even low concentrations (<1%) give rise to bulk refractive index responses that dwarf the actual binding signal of small molecules. Consequently, it is necessary to add DMSO to the running buffer before running the experiment in order to reduce this bulk refractive index offset. Despite best efforts, a difference in sample and assay buffer refractive indices can exist and require solvent calibration to resolve low analyte binding signals. The DMSO page uses micro-calibration injections of DMSO that can be included in assay protocols to construct a calibration curve that is then used to correct the final data set in advance of curve fitting.

To construct a solvent calibration plot you must prepare at least three solvent standards (and preferably six) that encompass the potential solvent concentration range of the sample buffer. For example, if an analyte
was injected across an immobilized ligand in a running buffer (PBS-T) containing 3% DMSO then a suitable six-point DMSO calibration curve might contain the standards 2.5, 2.7, 2.9, 3.2, 3.4, and 3.7% DMSO in PBS-T. These DMSO standards are then injected similarly to standard analyte samples in the same assay and their values then used to construct a calibration plot using the following procedure.

![Table Image]

**Figure 3-8: RI Calibration Step**

Next, the concentrations of the DMSO standard samples are set to the value ‘d’ to designate them as solvent standards (see “Correcting analyte concentrations” on page 7 for further information on reassigning and correcting concentration values). Next, the time limit lines are set in the RI Calibration step to construct the DMSO calibration plot. The before graph in the Prepare tab displays the time limit adjustment. The green and red lines are centered over a time point during the sample injection and the blue line is set directly on the injection stop.

![Graph Image]

**Figure 3-9: Time Limit Adjustment**

When the time limit lines are set properly, the **Configure** button can be pressed and a calibration plot is constructed when either Linear or Quadratic is selected in the pop-up window. The effect of varying solvent concentration is quantified and subtracted out of the sample signal. It is important to ensure that the samples lie within the solvent concentration range chosen, thus making this calibration appropriate.

**Blanks Step**

The composition of the running and sample buffer can have a large effect in SPR assays due to the presence of excipients. These can lead to an apparent change in the observed response even when injecting sample buffer in the absence of analyte (blank buffer injection). In order to generate the best quality data, it is important to double reference subtract that assay data. The preliminary reference subtraction is performed as shown in “Reference Step” on page 17 but to account for buffer effects the blank injection response must be subtracted from the entire data set.

The kinetic model fit will be inaccurate if this blank buffer injection response is ignored.

The random noise of a corrected curve is the sum of the noise from both working and reference curves. Therefore, a double-referenced response curve will possess more random noise than a single-referenced response curve.

Note that Octet® SPR Analysis automatically assumes that all samples with concentration equal to zero in the sam-
ple table are blank buffer injections. It is recommended to run at least three replicates of this blank and to choose the **Average** option in the Blank page (Figure 3-10). This will produce an averaged curve that is then subtracted from the full data set. Alternatively, selecting the **Closest** option will use only the curve that is nearest to the data set. Closest is more commonly used in large data sets run over many hours and where the buffer blank is repeated periodically.

![Figure 3-10: Blank Step](image)

Outlier buffer blank injections can be removed from the data set by double-clicking the outlier in the left graph and then clicking the **Remove** button.
Chapter 4:
Point Studies Tab

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Description of the Point Studies Tab

The Point Studies tab page enables rapid plotting of analyte dose response curves or other point-variable relationships. This tab contains a Point Selector which is optionally minimized at the top left side of the page.

A drop-down box allows different plots and analyses to be generated.

Affinity Plot

In an SPR assay, the analyte binding response should be proportional to the concentration of the analyte and therefore, dose response plots are a good test for outlier responses. In the absence of outliers, the user should expect a dose response curve similar to Figure 4-1. The dose response curve is obtained by positioning the $R_{eq}$ line (blue line in the Point Selector graph) on the response curve just before the end of the sample injection phase. The mean response of the Window size data region is calculated for each curve and plotted as a function of the analyte concentration. Readjustment of the position of the $R_{eq}$ lines will result in an immediate adjustment in the dose response curve.

Figure 4-1: Affinity Plot

Affinity plot: OneStep®

In addition to standard multi-cycle kinetics, plots can also be prepared for an analyte dose response curve for OneStep® injections. The number of $R_{eq}$ points can be adjusted with the up and down arrows. Figure 4-2 shows such a plot generated from multiple OneStep® injections of Trastuzumab binding erbB-2 (Her-2/neu).
NOTE: It is important to remember that equilibrium analysis of SPR data requires that each dose reach steady state, or an inaccurate affinity measurement will be made. More discussion on the specifics of model fitting in this page are shown in the following sections.
Affinity plot: formatting

All graph properties can be adjusted as desired. Right-click the graph and select Properties from the pop-up menu and alter parameters as required (Figure 4-3).

![Figure 4-3: Formatting the Affinity Plot](image)

Affinity plot: model fitting

As noted above, use of the Affinity model when analyzing dose responses necessitates all analyte concentrations have reached steady state. Note that dose responses are divided by flow channel with a new row of tabs, whereas the Point Selector graph will show all available flow channel responses. Checking the Fit check box (top left corner) will automatically fit a simple 1:1 steady-state affinity model to the data (Affinity Plot: Fit Check Box). The Fit check box to
the right of the $R_{\text{max}}$ parameter box designates $R_{\text{max}}$ as a fitted parameter in the model. Alternatively, if left unchecked, the $R_{\text{max}}$ value can be entered as a fixed value.

Figure 4-4: Affinity Plot: Fit Check Box
When multiple analytes are fit with the Affinity model, the Analyte MW will often require local $R_{\text{max}}$ values per Analyte. To enable a locally fitted $R_{\text{max}}$ for each analyte, click the L column heading in the Affinity table (Figure 4-5). The L column indicates the analyte $R_{\text{max}}$ is fit locally if checked, and the F column indicates the analyte $R_{\text{max}}$ is fitted rather than manually entered.

The prior examples show much better fits when local $R_{\text{max}}$ is allowed, because analyte MW varies by ~2-fold. The Show as % max allows the Y-axis of the affinity plot to be displayed as a response relative to the $R_{\text{max}}$ (Figure 4-6).
If a local $R_{\text{max}}$ is not desirable the data can be MW Corrected as outlined in “Entering and Correcting Analyte Concentrations and Names” on page 7 and the data can be fit with a global $R_{\text{max}}$ (Figure 4-7).
After the affinity model fitting, the $K_D$ is displayed in the Affinity table with standard error from the model fit. The desired representation of standard error can be changed using the Std. Dev. Display option located under the Configure menu.

The Affinity model can also fit a two-site equation assuming the analyte binds to two sites on the immobilized target, each characterized by a different binding affinity. To fit a two-site model click the box designated 2 in the sample table and a second site is introduced for all analytes. Individual analytes can be checked in this column as well.

Another data column will appear in the sample table when the check box Calc % Bound is selected. This option allows estimation of the percentage of analyte that will be bound for different mole ratios of analyte and ligand from the affinity constant returned in the model fit. Two parameter boxes appear to the right of the Calc % Bound check box allowing the concentration of drug (analyte) or protein (Target) to be entered. The % Bound displayed in the table is calculated for quick reference. See the Appendices for the equations used in the Affinity model(s) and the Protein Binding model.

**Hit Selection**

The purpose of the Hit selection feature is to enable rapid and accurate identification of fragment hits in the setting of a primary screen assay.

The necessary elements for Hit selection are: injections of fragments at sufficient concentration to elicit response (typically 100 - 500 µM), a positive control analyte at a near-saturating concentration injected periodically, a negative control analyte (optional) injected periodically, at least 96 fragment analytes and preferably 384 or more. The Hit selection processes primary SPR screening data by applying a LOESS (LOcal regrESSion) correction to the positive control and response data to account for parameters including drift, activity decay, and aberrations.

Using the Point Selector window, drag the blue line for one $R_{eq}$ point at the plateau of the injection. Figure 4-8 shows an example OneStep® data set.

![Hit Selection](image-url)
Similar to the Affinity plot, the active flow channels are separated by tabs in the Hit selection plot. Identify the positive and negative controls for the target immobilized on that flow channel using the drop-down boxes. Figure 4-9 shows the uncorrected Hit selection plot for an example screen.

Figure 4-9: Hit Selection Plot
The slide bars can be used to adjust the span settings for the control (positive control) and response LOESS corrections, respectively. Ideally, the positive control will be shown with a straight line across 100%. However, it is important not to choose the lowest span in order to avoid over-correcting the data. Figure 4-10 shows unacceptable LOESS span settings on the example screen.

![Figure 4-10: Unacceptable LOESS Span Settings](image)

The **Selection Method** can be chosen from: Box-Whisker, Norm Exp, or Combined. Box-Whisker is adjusted with the Hit Magnitude, Inter Quartile Range (IQR) Scale where a value of 3.0 is considered appropriate for most data sets. Lower IQR values will be less stringent when identifying hits and higher values will be more stringent. Norm Exp (normal exponential) is adjusted with the Hit Percentile Cut-off which defaults to 99.990, an appropriate setting for most data sets. Larger percentile cut-offs will be more selective than lower cut-off values. The Combined selection method enables both of the methods listed above and acknowledges overlap between hit identification with a "MultiHit" designation, where Hit designation means selection by a single method as opposed to both methods.

When a Hit selection method has been finalized, the Kinetics tab will show only hits and controls determined by that method. In this way the user can focus on kinetic and affinity analysis of the hits from the screen and ignore the non-binding analytes. In order to show all analytes in Kinetics, simply return the Hit selection drop-down menu to either None or Affinity.
# Chapter 5:
**Kinetics Tab**

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Kinetics Tab Introduction

The objective of the Kinetics tab is to allow kinetic modelling by non-linear regression curve fitting in order to extract the kinetic rate constants for the interaction. Figure 5-2 shows a typical kinetic page before curve fitting.

Figure 5-1: Kinetics Page

There are several elements to this page and the screen shot below indicates eight features that will be described.
1. **Response window:** The black curves illustrate the binding data to be fitted with the chosen model. The green and red time lines represent the injection start and injection end time points, respectively. These must be accurately positioned for successful curve fitting.

2. **Model selection:** The drop-down box includes an option to avoid fitting any curve (i.e. None). With Sites (#3 in Figure 5-2) set to 1, all models assume that the data can be approximated as a pseudo-first-order 1:1 interaction. Refer to Appendix D for a detailed discussion of each model.

   Selecting Simple off (i.e. $k_d$) allows only the dissociation rate to be fitted to the dissociation phase of the response curves.

   The Simple model (i.e. $k_a, k_d$) allows both the association rate constant and the dissociation rate constant to be determined.

   A two compartment model (i.e. Transport $k_{m}, k_a, k_d$) is provided for kinetic analysis where partial mass transport limitation is suspected. This model assumes the formation of an analyte gradient at the interface between bulk and surface thereby compensating for analyte transport limitation. Other special-case binding models are available and are discussed later.

   The ResSD value is the residual standard deviation of the fitted curves from the actual data curves.

   The Affinity (K) model fits a real-time isotherm to the data assuming all points are at equilibrium.

   **NOTE:** This is a reasonable assumption when the $k_d$ is 0.05 sec$^{-1}$ or faster.
3. **Sites selection:** This drop-down box allows the model to fit for multiple independent binding sites. This should be kept at 1 for most data sets. Other effects such as steric hindrance or rebinding may have a better fit with a 2-site model; however, this of itself does not confirm that two independent binding sites are present. Careful consideration of predicted biological function should be used when determining the number of sites to use with the model fit.

4. **Fitted parameter table:** The table includes the starting parameter values used to generate the simulated curves. When the fitting is complete these values will represent the optimized fitted parameters. The curves simulated from this initial value produce a set of simulated curves that differ from the actual experimental curves. Further fitting will result in simulated curves that differ very little from the actual experimental curves.

5. **Simulation graph:** Simulated response curves generated using the initial parameter estimates from the table and the simple association rate and dissociation rate model (i.e. $k_a$ and $k_d$). Prior to fitting, these simulated curves (shown in red) are different to the actual experimental data. After fitting is complete the fitted curves should superimpose upon the experimental curves.

6. **Residuals:** Residual plot showing the residual difference between the experimental curves and the simulated curves at each time point.

7. **Fit Button:** The Fit button begins the non-linear regression algorithm resulting in optimization of parameter estimates. The procedure performs successive iterations where the model parameters are changed in a systematic way in order to minimize the residuals.

8. **Kinetic Options Button:** The Options button provides a means of changing the model fitting in a variety of ways to suit your data set. See sections “Bivalent Analyte Model” through “Kinetics Options: Other Model Fitting Options” for more information on these fitting options.

9. **Estimate Button:** The Estimate button will provide initial estimates for any fitted parameter and can be clicked when a specific cell or cell range is highlighted in the Fitted parameter table. In some cases, the existing parameter value may need to be cleared before a new Estimate can be generated.
Kinetics: Dissociation Rate Constant

In some cases, the dissociation rate constant is fitted independently, such as when analyte concentration is unknown. The program fits the simulated model curves to the dissociation data to determine best-fit $k_d$ for each analyte. To identify the beginning of the dissociation phase drag the red line to the start of the dissociation phase in the data set. Click the Fit button to begin the least squares regression.

Figure 5-3: Dissociation Rate Analysis

Dissociation rate analysis is useful for off-rate ranking/screening and can be used to produce Avidity plots.
Kinetics: Association/Dissociation Rate Constants

The simple 1:1 model simultaneously fits the rate constants of association and dissociation and estimates binding affinity. Drag the green line to the injection start (often 0 sec) and the red line to the injection end.

**TIP:** Zoom into the data set around injection start and stop to accurately identify these two time points.

![Figure 5-4: Kinetics: Association and Dissociation Rate Constants](image)

After clicking the **Fit** button, the program begins the least squares optimization of the simulated curves. It is very important to pay attention to the placement of the vertical time lines that designate the start and end points for the association phase of the curve. Figure 5-5 shows the result of the model fitting procedure.
The simulated curves almost completely superimpose on the experimental data as shown by the low residual values, indicating a good model fit.

**Kinetics: Mass Transport Limited Kinetics**

Fitting the Transport \( (k_m, k_a, k_d) \) model is performed similarly where the mass transport rate constant is included in the model (Figure 5-6).

When fitting this model, it is important to note that fitted values for \( k_m \) of \( 1e^8 \text{ m}^3\text{s}^{-1} \) and greater represent no significant mass transport limitation in the majority of interactions. In this case the Simple \( (k_a, k_d) \) should be used.

If some analytes demonstrate transport limitation while others do not, the transport rate \( k_m \) can be fixed at \( 1e^9 \text{ m}^3\text{s}^{-1} \) which ensures mass transport is not limiting for the observed kinetic rates. Also, fitted values of \( k_m \) of \( 5e^5 \text{ m}^3\text{s}^{-1} \) or slower should be carefully considered as these are irregular values for the Octet® SF3 flow cells.

Figure 5-5: Curves After Least Squares Optimization
**TIP:** Mass transport limited kinetics are often characterized by linear association and slightly biphasic dissociation. Non-transport limited kinetics will have more rounded curvature in association and monotonic dissociation curves.

**Kinetics: OneStep® Injections**

OneStep® data analysis is similar to the analysis of conventional data sets with one main difference. In contrast to fitting conventional binding interaction curves, the concentration of analyte changes continuously during the injection and versions of the simple model and the complex models have been adapted accordingly. Figure 5-7 shows a OneStep® gradient assay in the kinetic tab of Octet® SPR Analysis. Note that the green solid line represents the relative analyte gradient concentration versus time.
Additional parameters appear on the Kinetics table when fitting OneStep® data. The analyte diffusion coefficient is a critical component of the analysis of OneStep® data because it defines the shape of the concentration gradient.

In addition to the previously discussed parameters there are two new factors shown in the Kinetics table. The $D_{app}$ is the apparent Diffusion Coefficient and the $N_{agg}$ is the Aggregation Number. The $D_{app}$ defaults to the value determined from the analyte molecular weight. In this case $N_{agg}$ will be at 1, indicating the analyte is a monomer of its ideal MW. Both of these can be fitted by selecting the $D_{app}$ column and floating these values. Then click **Fit** on the Kinetics Page. As the model fits $D_{app}$ to conform to the data it will adjust the $N_{agg}$ respectively.

Refer to see “Kinetics Options: OneStep® Injection Options” on page 46 for other specific model fitting considerations.

**Kinetics: NeXtStep™ Injections**

NeXtStep™, like OneStep®, enables the injection of concentration gradients of analyte(s) which can be fit with standard kinetic and affinity models. When a NeXtStep™ injection is used for direct analyte binding (not in a Competition or Inhibition method), the resulting data can be fit with the Simple ($k_a, k_d$), Transport ($k_m, k_a, k_d$), Affinity ($K_D$), or other available binding models.

A key difference of NeXtStep™, compared to OneStep®, when analyzing binding data is that diffusion is not a parameter in the model fit since the gradient is not defined by analyte diffusion theory. Rather, analyte dispersion is empirically observed using a surrogate sample (typically Sucrose) and the gradient is defined by concentration coefficients. Methods for analyzing competition or inhibition data using NeXtStep™ will be addressed in specific Technical Notes.
Affinity Model

A real-time equilibrium isotherm model (Affinity \(K_D\)) can be applied to data with sufficiently fast dissociation rate constants (0.5 sec\(^{-1}\) or faster). This model assumes that all points are at steady state and fits for equilibrium dissociation constant \(K_D\) and \(R_{\text{max}}\) only. Figure 5-8 shows an example of an Affinity isotherm fit with a two-site interaction.

Bivalent Analyte Model

The Bivalent Analyte Model represents data where the analyte has two (or more) identical binding sites which can bind the immobilized Ligand individually or simultaneously. The following is the rate equation describing the Bivalent Analyte model:

\[
A + 2B \rightleftharpoons AB + B \rightleftharpoons ABB
\]

In the equation, \(B\) is Ligand and \(A\) is Analyte which can bind two Ligand molecules. The most common example of Bivalent Analyte is an IgG antibody molecule analyte binding immobilized antigen. In this example, each antibody binding site can bind antigen and after the first site binds antigen (Ligand), the second site is able to bind a neighboring antigen at a different (due to proximity) rate. Since the Bivalent Analyte model is more complex, it is recommended that the user read Appendix A for guidelines on when to apply a more complex kinetic model but it in general it is recommended to design assays so that the simplest model (1:1) can be assessed.
Note that $k_{a2}$ is in terms of RU$^{-1}s^{-1}$ and requires conversion by refractive index increment to be interpreted as M$^{-1}s^{-1}$.

An in-depth review of the Bivalent Analyte model is shown in the work by MA Cooper & DH Williams (1999)\(^1\), where SPR was used to compare the kinetics of antibody binding antigen (ligand) and antigen binding antibody (ligand).

### Two-State Model

The Two-State Model represents data where the analyte binds ligand forming reversible complex (AB) which can proceed to a second state (AB'). The second state may be a different conformation of ligand or complex, covalent reaction, etc. The rate equation describing the Two-State model is shown in Figure 5-12.

$$
\begin{align*}
A + B & \rightleftharpoons AB \rightleftharpoons AB' \\
& \;
\end{align*}
$$

![Figure 5-10: Two-State Model Equation](image)

In the equation, B is Ligand, A is Analyte, AB is the initial bound complex, and AB' is the bound complex in its second state. Since the Two-State model is more complex, it is recommended that the user read Appendix A for guidelines on when to apply a more complex kinetic model.

### Two-State Irreversible Model

The Two-State Irreversible model is a simplification of the Two-State model, wherein complex that has changed to the second state AB' does not readily return to the initial state AB. In the above rate equation, parameter $k_{d2}$ is a fixed constant of 0 sec$^{-1}$. When this model is selected, an additional term $C_c$ - the commitment to covalency - will be displayed in the Kinetics result table. $C_c$ is defined by the following relationship:

$$
C_c = \frac{k_{a2}}{k_{a2} + k_{d1}}
$$

![Figure 5-11: Two-State Irreversible Model Equation](image)

It is recommended that the user read Application Note 23, *Commitment to covalency: kinetics of irreversible inhibitors with a regenerable streptavidin sensor on the Pioneer FE system* for a complete set of guidelines on when to apply a more complex kinetic model such as the two-state irreversible model.

---

Diffusion Model

The Diffusion model can be used to fit the apparent diffusion coefficient of an analyte using only the refractive index signal when a diffusion-based injection (i.e. OneStep®) is used. This model assumes a linear relationship between concentration and SPR response exists (i.e. non-binding refractive index response). Therefore, this model should not be used for interactions where kinetic or affinity binding is occurring. For most biomolecules, a concentration of >0.1 mg/mL would be necessary to observe a reliable response. Concentrations of >100 mg/mL analyte can be tested unless significant viscosity or precipitation is issue. Figure 5-12 shows a OneStep® Pulse injection of a 3% sucrose solution fitted to the Diffusion model.

The fitted parameters for the Diffusion model are $R_{\text{max}}$ (RU) and $D_{\text{app}}$ (m$^2$/s). $R_{\text{max}}$ represents the signal at maximum concentration, which is observed for sigmoidal OneStep® but is not observed for OneStep® Pulse. Since refractive index is expected to be linear to analyte concentration, $R_{\text{max}}$ in this context can be explained as:

$$R_{\text{max}} = C \cdot \eta_d$$

where $C$ is analyte concentration (typically in mg/mL) and $\eta_d$ is refractive index increment (RU*mL/mg). Alternative units can be used where available. This can be viewed as a QC metric for the sample response which should give an $R_{\text{max}}$ value within the correct range for known concentration and $\eta_d$.

Aggregation Model

The Aggregation model can be used to quantify the relative concentrations and diffusion coefficients of components in a bi- or poly-disperse sample using only refractive index signal from a diffusion-based injection (i.e. OneStep®). The basic assumption with the Aggregation model is that the sample comprises at least two analyte components that differ by diffusion coefficient (i.e. monomer and heptamer components, etc.). A requirement for the Aggregation model is that the total analyte concentration and the analyte refractive index increment ($n_d$ or $dn/dc$).
be known. Fitted parameters of the Aggregation model are refractive index increment of monomer (dndc_m), diffusion coefficient of monomer (D_m), concentration of aggregate species (C_a), refractive index increment of aggregate species (dndc_a), diffusion coefficient of aggregate species (D_a).

Output parameters from the Aggregation model are concentration of monomer species (C_m) and percent aggregate (%agg). Ideally dndc_m and to a lesser extent, dndc_a, should be known. Dndc for proteins >10 kDa can typically be assumed to be 188.8 RU*mL/mg and predictions for a wide panel of proteins have shown a range of 180 – 200 RU*mL/mg\(^1\). A wider distribution is seen for smaller proteins, but the mean dndc is still similar at 190.2 RU*mL/mg. Note that the Aggregation model enables a separate Retention term for monomer (RF_m) and aggregate (RF_a). Refer to “Retention” on page 47 for more information.

Kinetics Options: General Options

The Kinetics Options button accesses additional parameters for modelling complex data or data with specific artifacts. Kinetics options can be selected by clicking the Opts button as shown in position 8 in Figure 5-2. The top tool bar has four buttons; the Fix button is used to designate parameter values as constant for a model fit. By default, all parameters in the Kinetics Options pop-up window are fixed unless they are manually floated. The Float button designates parameter values as variables that must be iteratively fit.

Figure 5-13: Kinetics Options

Link tab

The Kinetics Options pop-up allows each fitted parameter to be fitted as a local parameter or as a global parameter. In Octet® SPR Analysis, this local/global indication is called Link. The parameters of any particular model may be constants or variables and may be fitted separately to each curve in the set representing various analyte concentrations (i.e. local fitting) or constrained to the same value for the set of curves (global fitting). Parameters should be constrained when this is possible as it results in a more robust fit. However, there are occasions where it is not correct.

to fit all parameters globally. For example, in affinity capture (a common kinetic assay format), the analyte binding capacity (i.e. \( R_{\text{max}} \)) is not constant for all interaction curves because there is experimental variability during the ligand capture step. This variation in capacity may be dealt with by fitting the \( R_{\text{max}} \) as a local variable. To do this click the \textbf{Link} tab in the Kinetics Options. Figure 5-13 shows this window. Each row in the table represents an individual curve. During the curve fitting process, the fitting algorithm fits the model parameters to the first curve in the table. If "1" is entered in all rows for all other curves, the algorithm assumes that there is only one value for this parameter for the data set. Note that 1 corresponds with the injection ID number in the far-left column thus linking all injections to injection #1. Figure 5-13 shows all other curve parameters are linked to injection #1 which ensures that the \( k_a \), \( k_d \) and \( R_{\text{max}} \) are all fitted as global parameters (i.e. only a single value is allowed for the entire data set). However, if the values in the \( R_{\text{max}} \) column are deleted as shown in Figure 5-14, \( R_{\text{max}} \) will be fitted locally (per curve). After adjusting this table, press OK and press the Fit button. The different \( R_{\text{max}} \) values for each curve are now shown in the Kinetics page parameters table.

![Figure 5-14: Link Table](image)

Similarly, it is also possible to fit \( k_a \), \( k_d \) and \( k_m \) as local parameters but this is not considered best practice and assay optimization should be prioritized.

**Begin and End Tabs**

The Begin and End pages serve the same function as the vertical time lines on the plot. They simply designate the start and stop times for the association phase of an interaction curve. Usually a set of interaction curves will possess the same start and stop time points and the vertical time lines are a useful way of conveniently setting these for the model. This corresponds with the All radio button selection. However, there are non-ideal situations where it may be desired to fit a model to a set of data with association phases of different durations. In this case, go to the \textbf{Begin} page and select \textbf{Separate} as shown in the screen shot on the right in Figure 5-15. The actual start time for each association phase is typed into the table for each curve. The values appear red indicating that they are constants and will be fitted as such. Highlighting these values and pressing the \textbf{Float} button will designate these values as local parameters.
to be fitted but this is not recommended. The end time points may be added in a similar fashion in the End page. If the injections are set to the same start/stop times (All), the Fit check box allows the model fitting to find the best fit for these values as well.

Figure 5-15: Kinetic Options: Begin Tab

Step RI Tab
Most of the data analyzed in this Tutorial does not possess significant bulk index offsets and it is unusual to have such signal offsets when reference curve subtraction is employed. However, it is possible to observe bulk index offsets, especially when a large differential in mass loading exists between the working and reference flow channels. This can result in the excluded volume effect where volume excluded by protein modulates the sensitivity of the optical channel for changes in refractive index. These appear as steps at the beginning and end of the injection (i.e. association phase) and are due to bulk refractive index variations or in some cases bulk refractive index offsets due to an excluded volume effect. The simulated curve shown in Figure 5-16 shows a bulk index offset.

Figure 5-16: Curve with Index Offset
Octet® SPR Analysis can add steps to the simulated curves to account for these offsets. To use this function, press the Options button on the Kinetics tab page tool bar. The pop-up in Figure 5-17 will appear.

The Step Refractive Index (Step RI) page is shown in the following two screen shots. The screen shot on the left shows that the bulk refractive index value for each curve is set at zero and is designated a constant. If the data includes bulk refractive index offsets, these will need to be fitted locally to each curve during model fitting. To do this, simply highlight the column of values and then click the **Float** button. The values will appear black. Click **OK**. The pop-up will close and the program will return to the Octet® SPR Analysis kinetics page. Select the **Fit** button. The model is then fitted to the data set but the bulk refractive index offsets are included in the fit. After fitting, re-open the Kinetic options pop-up and the optimized response steps for each curve are displayed as shown on the right in Figure 5-17.

**NOTE:** Step RI can be fit to OneStep® injections as well, where the RI shift is applied through the entire association phase regardless of gradient profile.

This function allows accurate model fitting when bulk refractive index offsets exist in the data.

### Kinetics Options: OneStep® Injection Options

#### Analyte RI

Analyte RI refers to a refractive index offset in the data attributed to the inherent refractive index of the analyte. In OneStep® injections, the analyte concentration changes versus time, so this RI term adjusts according to the analyte concentration profile. This term can also be used in many cases to model refractive index offsets due to buffer solvents such as DMSO.

**NOTE:** The Solvent RI term is also available to model solvent RI offsets.

Care should be taken during sample preparation to minimize refractive index offsets as described in “RI Calibration Step” on page 18. This term can be floated to remove these effects from referenced data for conventional and gradient injection types.
Solvent Diff & Solvent RI

Solvent Diff and Solvent RI are two corresponding parameters that model solvent RI offsets specifically in OneStep® data. Solvent Diff (diffusion coefficient) represents the diffusion coefficient of the solvent offset or discrepancy. This is a value which requires manual input and cannot be fit as a floated parameter. A fast diffusion coefficient often describes this offset behavior (for example 1e⁻⁹ to 5e⁻⁹). Solvent RI will not function without a manually input Solvent Diff coefficient. However, once Solvent Diff is input, Solvent RI can be fit as a floated parameter. As described in “RI Calibration Step” on page 18, the solvent offsets in a sample should be minimized, if possible, and Solvent RI is intended only for those which may remain after calibration.

Retention

Retention in Kinetics Options refers to analyte retention in the capillary tube during a OneStep® injection. The retention term can be fitted if there is concern that the analyte is interacting with the tubing as it moves towards the flow cell. This will show in the data as a curve that binds later than expected, and so the model will be shifted to the right in time. This can be fitted by selecting the Retention values in red and clicking on the Float button at the top. A local fit for each analyte can be done by removing the numbers in the Link column; however, this is not as robust as the recommended global fit routine.

The retention term is a value between 0 (no analyte retention) and 1 (complete analyte retention). For best results, ensure that retention does not exceed 0.3 as this may indicate that a strong hydrophobic event is occurring between analyte and tubing. In some cases, retention can be fixed at a value greater than 0.3 but the analysis is not as robust if retention is present at such a high level. Often buffer conditions must be modified to prevent analyte adsorption/ sticking to tubing or other surfaces. Figure 5-19 shows a simulation of moderate affinity binding ($K_D \sim 600$ nM) with different retention factors where 0.3 retention factor represents a significantly retained interaction.
The final tab in the Kinetics Options pop-up is the Options tab. The Rate upper limit box specifies the maximum allowable \( k_a \) for a model fit and improves the reliability of the fitting procedure by bounding this parameter. The end Chisq (i.e. \( c^2 \)) change defines the cut-off point at which the difference between the simulated and experimental curves is considered minimal; the optimization procedure is terminated when this value is reached. The Max number of iterations box sets the upper limit on the number of iterations the curve fitting algorithm will run before stopping. The Update Display option sets the frequency for updating the fitted graphs. For example, a value of 1 will update the fitted graphs after every iteration of the curve fitting algorithm. A larger value will update the graph less frequently during curve fitting but may result in faster fitting as time is not wasted updating the graph to show intermediate results. Check Display as % \( R_{\text{max}} \) if you would like your y axis displayed in this manner. Show Zero Conc Curves will include the subtracted blanks in the plot. The Refit options will cause the fitting algorithm to restart if it cannot get below the input restrictions.
Ligand efficiency (LE) as a concept is a means of comparing molecules by their binding energy per atom. Therefore, the Octet® Discovery software will collect heavy atom count (HAC) at the protocol design stage for use in calculating ligand efficiency in Octet® SPR Analysis. The equation below defines the relationship between HAC and $K_D$ to estimate ligand efficiency\(^1\).

$$LE = \left(-2.303 \times \left(\frac{RT}{HAC}\right)\right) \times \log K_D$$

R is the ideal gas constant, T is the analysis temperature in Kelvin, HAC is the integer value of non-hydrogen (heavy) atoms in the molecule and $K_D$ is the equilibrium dissociation constant. The ligand efficiency metric can be enabled throughout the Octet® SPR Analysis workflow by selecting it from the Configure > Advanced Metrics menu.

---

Once enabled, a new LE column will appear in the Kinetics grid next to the $K_D$ column. Ligand Efficiency can also be displayed in Results graphs by right-clicking the graph, selecting Properties, and configuring the LE option under the Inset options as shown below. $K_D$ is required to be in the inset table before LE can be enabled.
Results Tab Introduction

The Results tab displays the plots generated during Octet® SPR Analysis analysis. Plots can be sorted by analyte or ligand using the respective buttons in the upper tool bar. Views of the time data or dose responses can be viewed by clicking the respective buttons on the lower level of the upper tool bar. The Layout drop-down menu gives the graph tiling options and when the visible graphs exceeds the Layout setting, the Page tab buttons allow each Page to be scrolled through.

Right clicking over a particular plot and clicking Properties will allow that plot to be formatted. When the plot is ready it can be exported as a graph, or as a raw data set (i.e. text file) by right clicking and selecting Copy graph or Copy data, respectively. Additionally, Configure in the file menu has the Copy visible graphs option which will copy an image of the current Layout view to the clipboard.

![Figure 6-1: Results Page](image)

Use the Hide button on the main toolbar to hide any undesired plots or the Filter drop-down boxes to isolate specific plots.

Clicking the Export button on the upper tool bar will allow the export of data in two text file formats and one image file format (.png). Clamp data files (.txt) include the simulation curves from model fitting if replotting in another software is desired. Image files will be created for all the selected views. This is a simple method to produce numerous image files from screening data or other large data set.
Chapter 7:
File Saving
File Saving

Use the **Save** button on the main tool bar to save analyzed data. Saved Octet® SPR Analysis files appear as .qdt files and raw data files from the Octet® SPR Discovery software are exported as .qdtx files.

![Save Button](image1)

**Figure 7-1:** Save Button

The Print pop-up (*File > Print*) has a variety of options to choose from, as shown in Figure 7-2.

The Experiment tab presents a text window where one may add notes to the data set.

![Print Screen](image2)

**Figure 7-2:** Print Screen
Appendix A:
Introduction to Non-linear Regression

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- Assessing the Goodness of Fit ............................................................. 56
- Is the Model Appropriate? ................................................................. 56
Description of Non-linear Regression

Octet® SPR Analysis software employs non-linear regression and the Levenberg-Marquardt algorithm to fit experimental data to a binding interaction model that defines the interaction. The parameters of any particular model may be constants or variables and may be fitted separately to each curve in a set representing various analyte concentrations (i.e. local fitting) or constrained to the same value for the set of curves (global fitting). The Octet® SPR Analysis software fits the model to the entire data set as opposed to defining subsets of data for association and dissociation phases. This can be considered as a global time constraint and further improves the reliability and robustness of the fit. Global analysis of the data set does require very high data quality, but both Octet® SPR Discovery and Octet® SPR Analysis software have been optimized to achieve this.

Initial Parameter Estimates

Before fitting a model, initial parameter values must be entered that are reasonable values. Usually initial parameter estimates that are orders of magnitude distant from the true value do not cause any problems. When activated, the program iteratively changes the fitted parameters such that the average residual difference between the simulated curves and the experimental curves is minimized. The program stops when further reduction of the residuals becomes insignificant. However, it is best to choose values that are reasonable estimates as the model fitting algorithm may sometimes fail to find a unique solution and get trapped in a local minimum (i.e. a false minimum that is not the true minimum). This is an unavoidable property of all curve fitting programs. It is also more common when fitting complex models with many parameters. To avoid this problem, simply observe the simulated curves before initiating a fit and manually iterate the initial parameter estimates until these simulated curves roughly match the actual experimental data.

Assessing the Goodness of Fit

The goodness of fit is measured as the averaged residual standard deviation. Usually, if the averaged standard deviation of the residuals is less than 1% of the maximum response for the curve set, this indicates an excellent fit. However, values that exceed 10% are sometimes acceptable (e.g. very low binding responses i.e. <20 RU). It is important to remember that the standard deviation of the residuals can never be lower than the standard deviation of the random baseline noise response for the instrument.

Is the Model Appropriate?

It is important to always consider the biological and physical relevance of any particular model to a data set. A general recommendation is to design the assay such that the simplest model can be used for assessment and begin fitting using the simplest model – a reasonable fit to this model is more likely to return good parameter estimates than more complex models that fit with lower residuals. For example, the two-compartment model (Transport \((k_{m}, k_{a}, k_{d})\)) may produce lower residuals than when the simple model is fitted, but is this appropriate for your data? If the observed binding rate of your interaction is relatively slow (i.e. \(<1e5 M^{-1}s^{-1}\)) and the \(R_{max}\) is limited appropriately (i.e. \(<200 RU\)), then the two-compartment model is not likely required. Often experimental artifacts can give rise to complex or heterogeneous binding curves which do not reflect a complexity in the interaction itself, but rather complexity in the experimental design or sample condition. Fitting more complex models can be compelling when heterogeneous data sets require explanation, but there is danger in over-misinterpreting the data set on the basis of an unsupported complex model fit. It is better to design experiments where the pseudo-first-order model is a reasonable approximation than to fit more complex models to your data.
Appendix B:
Defining the Expected Upper Limit for the Association Rate Constant

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Defining the Expected Upper Limit for the Association Rate Constant

When fitting a model there are a number of checks one can perform to assess the consistency of the parameters that are returned. For example, is it reasonable if an association rate constant of \(10^9 \text{ M}^{-1}\text{s}^{-1}\) is returned for an interaction of a 100 kDa analyte with an immobilized antibody? We can answer this question by understanding the biological and physical meaning of the association rate constant.

For the vast majority of interactions, the on-rate \((k_a)\) cannot exceed a rate that is limited by molecular diffusion. This diffusion limitation is not related to the mass transport limitation discussed previously with reference to the two-compartment model but to the intrinsic rate of binding. For example, increasing the viscosity will lower the \(k_a\) and increasing the temperature will increase the \(k_a\). This is predicted from the Smoluchowski equation. The following simulation assumes:

- Temperature = 25 ºC
- Viscosity and salt = standard buffer conditions (i.e. 0.15 M salts, pH 7.4)
- Ligand has a diffusion rate of zero
- Ligand MW = 150,000 Da
- Ligand Gyration radius = \(3.6 \times 10^{-9} \text{ m}\)

A modified Smoluchowski equation can be used to estimate the maximum possible on-rate \((k_{max})\).

\[
k_{max} = 4000 \times \pi \times r_{ab} \times K \times f \times N_A \times (D_A + D_B)
\]

Figure B-1: Modified Smoluchowski Equation

where

- \(r_{ab}\) = the sum of the gyration radius for the ligand and analyte
- \(K = 0.05\) = interaction constant related to binding area, i.e. 25% area of A multiplied by 20% area of B thus \(0.25 \times 0.2 = 0.05\).
- \(f = 1\) = assumes that electrostatic forces are not dominant
- \(N_A = \text{Avogadro's number} = 6.023 \times 10^{23} \text{ mol}^{-1}\)
- \(D_A, D_B\) = Diffusion coefficient for the analyte (A) and ligand (B) respectively. If B is immobilized then \(D_B\) will be zero
Defining the Expected Upper Limit for the Association Rate Constant

Figure B-2: Plot of Smoluchowski equation

Estimated variation in $k_{\text{max}}$ as a function of molecular weight under diffusion limited conditions (i.e. these represent the maximum possible $k_a$ values at each molecular weight). The active site area of the analyte was held constant at $1.8 \times 10^{-18} \text{ m}^2$.

The plot for this simulation clearly indicates that an analyte with a molecular weight of 100 kDa cannot be expected to produce an association rate constant of $10^9 \text{ M}^{-1}\text{s}^{-1}$ and an association rate constant $<3 \times 10^7 \text{ M}^{-1}\text{s}^{-1}$ is expected. Therefore, there must be an error in the model fit.

Interactions do exist where the maximum association rate exceeds the limit set by diffusion limitation but these are not common. For example, DNA-binding proteins may be electrostatically steered into the binding site by long range electrostatic forces\(^1\). Thus the electrostatic factor may be in the range 1-100. This mechanism may not be exclusive to DNA-protein interactions but it is not dominant for most affinity ligands. There are other factors at the sensing surface that may influence the $k_a$. Other spatial effects may become significant at a surface, such as the loss of one rotational degree of freedom, surface electrostatics, local pH and ionic strength variations at the surface, and the chemical aspects of immobilization.

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Appendix C:

Mass Transport

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Description of Mass Transport

In the following discussion it will be assumed that each flow cell possesses a height, width and length of 0.003 mm, 0.3 mm and 4 mm, respectively.

How does flow rate affect mass transport?

The mass transport limitation (MTL) factor was calculated under different flow rates keeping all other parameters constant to answer this question. A value of 1.0 indicates complete mass transport limitation and low values indicate kinetic limitation.

The SPR signal is an averaged signal over the interaction area surface. Transport of analyte to the surface is the result of convective and diffusion forces. These phenomena are described by the mass transfer coefficient \( L_m \) and are related to the flow cell design and operational flow rate. A good introduction to these principles can be found elsewhere\(^1\) but a brief review will be provided here. The delivery of analyte in the bulk \( (A_0) \) to the sensing surface \( (A) \) may be described by a mass transport coefficient:

\[
A_0 \rightleftharpoons A \quad \frac{k_m}{k_m}
\]

**Figure C-1: Mass Transport Coefficient**

where the forward and reverse rate constants are the mass transport coefficient, \( k_m \).

The interaction at the sensing surface is as follows.

\[
A + B \rightleftharpoons AB \quad \frac{k_a}{k_d}
\]

**Figure C-2: Interaction at Sensing Surface**

where A and B are the analyte and ligand, respectively; AB is the complex formed; and the forward (on) and reverse (off) rates are \( k_a \) and \( k_d \), respectively.

The first reaction describes transport of the analyte to the surface and the second reaction scheme describes consumption of the analyte at the surface via an affinity interaction with an immobilized ligand. In most experiments, it is not possible to attain 100% kinetic limitation nor is it possible to attain 100% mass transport limitation (MTL). How-

---

ever, by changing the concentration of immobilized ligand or the flow rate it is possible to control the mass transport characteristics. This is implied in the following relationship:

\[ MTL = \frac{L_r}{L_r + L_m} \]

**Figure C-3: Mass Transport Limitation Relationship**

where:

- \( L_r = K_a[B] \) = Coefficient of reaction flux
- \( L_m \) = Mass transport coefficient (see expression 4)
- \([B]\) = Immobilized ligand concentration

Hence, kinetic experiments require high flow rates and low ligand concentrations, whereas high MTL conditions require the opposite. Most applications will require kinetic limitation, but there are some interesting applications requiring high MTL conditions.

When using a flow cell, it is important to minimize the height of the unstirred diffusion layer that exists at the interaction surface. This can be accomplished by changing the geometry of the flow cell and the flow rate.

\[ h_{diff} \approx \frac{3}{F} \sqrt[3]{D \cdot h^2 \cdot b \cdot l} \]

**Figure C-4: Changing the Geometry of the Flow Cell and Flow Rate**

where:

- \( h_{diff} \) = Height of diffusion layer (m)
- \( D \) = Diffusion coefficient of the analyte (m\(^2\)s\(^{-1}\))
- \( h, b, l \) = Height, width and length of flow cell (m)
- \( F \) = Bulk flow rate (µL min\(^{-1}\))

We assume that the flow cell is 4.0 mm long, 0.3 mm wide, and 0.03 mm high, and a reasonable flow rate of 30 µl/min. For the detection of a 50 kDa analyte, from expression (3), we estimate \( h_{diff} \) to be 5.5 mm. These calculations require the diffusion coefficient for the analyte, which may be estimated from Stokes law and the Einstein-Sutherland equation:
where:

\[ k = \text{Boltzmann constant} = 1.381 \times 10^{-23} \text{ JK}^{-1} \]

\[ T = \text{Absolute temperature (K)} \]

\[ MW = \text{Analyte molecular weight (Da)} \]

\[ NA = \text{Avogadro's number} = 6.022 \times 10^{23} \text{ mol}^{-1} \]

\[ \eta = \text{Viscosity of liquid} = 0.001 \text{ Kg/m/s} \]

\[ f/f_0 = \text{Friction factor} = 1.2 \]

The SPR signal is an averaged signal over the interaction area. Transport of analyte to the surface results from convective and diffusion forces as discussed previously. These phenomena are described by the mass transfer coefficient \((L_m)\) and are related to the flow cell design and operational flow rate by:

\[ L_m = C_{lm} \sqrt[3]{\frac{D^2 \times F}{h^2 \times b \times l_2}} \]

where:

\[ C_{lm} = 1.47 \left( 1 - \left( \frac{l_1}{l_2} \right)^2 \right) \right) \]

\[ l_1 \text{ and } l_2 \text{ are the start and end of the active sensing region relative to the inlet position.} \]
What is the upper $k_a$ limit that can be estimated?

Myszka et al.\textsuperscript{1} (1999), demonstrated that the two-compartment model can resolve kinetic constants in the presence of 50% mass transport limitation. In general, medium to high molecular weight analytes (e.g. most proteins) will not have appreciable mass transport limitation. In fact, only very low molecular weight analytes will suffer mass transport limitation and only if capable of binding ligand at the maximum theoretical rate, which is unlikely.

Appendix D:

Pseudo-first-order Interaction Models

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Simple Kinetic Model \((k_a, k_d)\)

A biomolecular interaction between a soluble analyte \((A)\) and an immobilized ligand \((B)\) can be interpreted in terms of pseudo-first-order kinetics, where the rate of formation of complex is described by the following differential equation:

\[
\frac{dAB}{dt} = k_a A(t) B - k_d AB(t)
\]

\[
AB(0) = 0
\]

**Figure D-1**: Simple Kinetic Model

where:

- \(AB(t)\) = Molar concentration of complex at the interaction surface at time \(t\)
- \(A(t)\) = Molar concentration of analyte at the interaction surface at time \(t\)
- \(B\) = Molar concentration of available immobilized ligand
- \(k_a\) = Association rate constant \((\text{M}^{-1}\text{s}^{-1})\)
- \(k_d\) = Dissociation rate constant \((\text{s}^{-1})\)

Octet® SF3 instruments monitor surface interactions, and produce real-time interaction curves where the rate of change of response is directly related to the formation of complex AB. Equation 1 can be rewritten then in terms of response:

\[
\frac{dR}{dt} = k_a A(t) R_{max} - k_d R(t)
\]

\[
R(0) = 0
\]

**Figure D-2**: Kinetic Model in Response Terms

where:

- \(R(t)\) = Sensor chip response at time \(t\)
- \(R_{max}\) = Maximum response if all available ligand binding sites are occupied

This model holds for multi-cycle kinetics, as well as time-varying concentration injections. When employing flow cell-based analysis, the analyte concentration at the interaction surface can be approximated by the injected analyte concentration at any time, assuming mass transport of analyte to the surface is not limiting.
\[ A_0(t) = A(t) \]

Figure D-3: Kinetics Model with Flow Cell-Based Analysis

where:

\[ A_0(t) = \text{Molar concentration of injected analyte sample at time t} \]

\[ A(t) = \text{Molar concentration of analyte at the interaction surface at time t} \]

**Mass Transport Limited Kinetic Model \((k_m, k_a, k_d)\)**

The analysis outlined previously describes a simple biomolecular interaction model that is governed by 1:1 pseudo-first-order kinetic behavior. If mass transport limitation is expected to be significant then the \(k_m, k_a, k_d\) model should be fitted to the data set. In this model, the analyte concentration at the interaction surface is not approximated by the injected analyte concentration because mass transport of analyte to the surface is limiting. As a consequence, a second rate equation must be included to account for the analyte gradient at the sensing surface:

\[
\frac{dA}{dt} = k_m * (A_0 - A) - (k_a * A * B - k_d * AB)
\]

Figure D-4: Kinetic Model that Accounts for Analyte Gradient

This rate equation is integrated and coupled to equation 2, allowing the rate of formation of complex to be modeled when mass transport limitation is significant. This is also referred to as a two-compartment model.

**Analytical Solutions for Multi-Cycle Kinetics**

For multi-cycle kinetic injections, the analyte concentration from equation 2 is constant with respect to time, and an analytical solution exists. During the association phase, when \(A(t) = A_0\), the response at time \(t\) is given by:

\[
R(t) = \frac{k_a * A_0 * R_{max} \left(1 - e^{-(k_a * A_0 + k_d)(t-t_0)}\right)}{k_a * A_0 + k_d}
\]

Figure D-5: Fixed Concentration Model: Association Phase

where:

\[ t_0 < t < t_1 \]

\[ t_0 = \text{Injection start time} \]

\[ t_1 = \text{Injection end time} \]
During the dissociation phase, when \( A(t) = 0 \), the response at time \( t \) is given by:

\[
R(t) = R_0 e^{-k_d t}
\]

**Figure D-6: Fixed Concentration Model: Disassociation Phase**

where:

\[ t > t_1 \]

\[ R_0 \] = Sensor response at time \( t_1 \) (typically found by solving equation 5 at time \( t_1 \))

### Concentration Profiles for OneStep® Injections

Octet® SF3 instruments include novel injection types where the analyte concentration is not constant during the injection. For OneStep® injections, the concentration profile is given by the following equation:

\[
A_0(t) = \frac{1}{2} \left( 1 - \text{erf} \left( \frac{1 - \frac{t - t_0}{\tau}}{\sqrt{2 k(t - t_0) / vL\tau}} \right) \right)
\]

**Figure D-7: Concentration Profiles for OneStep® Injections**

where:

\[
k = (11 - 16r + 6r^2) \left( \frac{v^2 * d^2}{192D} + D \right)
\]

\[
\tau = \frac{\tau_0}{r} - t_0
\]

\[
\tau_0 = \frac{\pi L d^2}{4f}
\]

\[
v = r \frac{4f}{\pi d^2}
\]
The velocity through the dispersion loop:

\[ d = \text{Diameter of the dispersion loop (m)} \]

\[ L = \text{Length of the dispersion loop (m)} \]

\[ f = \text{Flow rate (m}^3\text{s}^{-1}) \]

\[ D = \text{Apparent diffusion coefficient of the analyte} \]

\[ r = \text{Retention factor (0 < r < 1) (unitless)} \]

The Diffusion model may be used to estimate the apparent diffusion coefficient by directly fitting the concentration profile to the SPR data when a measurement is made of the analyte’s bulk refractive index signal (no binding to immobilized ligand).

**Steady State Affinity Model**

Under steady state interaction conditions, the biosensor response varies with concentration according to the relationship:

\[ R(c) = \frac{c \times R_{\text{max}}}{K_D + c} \]

*Figure D-8: Steady State Affinity Model*

where:

\[ c = \text{Molar concentration} \]

\[ R_{\text{max}} = \text{maximum response is all available ligand binding sites are occupied (in RU)} \]

\[ K_D = \text{Equilibrium dissociation constant (in Molar concentration)} \]
Bivalent Analyte Model

A bivalent analyte is one that can bind to one or two ligand molecules. Binding is expressed using two sets of rate constants, the first describing binding of the analyte to a first ligand molecule, and the second describing the binding of an analyte-ligand pair to a second ligand molecule. The binding model is governed by the following set of differential equations:

\[
\frac{dB}{dt} = -\left( k_a \cdot A(t) \cdot B - k_d \cdot AB(t) \right) - \left( k_{a2} \cdot AB(t) \cdot B - k_{d2} \cdot AB2(t) \right)
\]

\[ B(0) = R_{max} \]

\[
\frac{dAB}{dt} = \left( k_a \cdot A(t) \cdot B - k_d \cdot AB(t) \right) - \left( k_{a2} \cdot AB(t) \cdot B - k_{d2} \cdot AB2(t) \right)
\]

\[ AB(0) = 0 \]

\[
\frac{dAB2}{dt} = k_{a2} \cdot AB(t) \cdot B - k_{d2} \cdot AB2(t)
\]

\[ AB2(0) = 0 \]

Figure D-9: Bivalent Analyte Model

where:

\( A(t) \) = Molar concentration of analyte at the interaction surface at time \( t \)

\( B \) = Available immobilized ligand

\( AB(t) \) = Concentration of complex (single analyte-ligand binding) at the interaction surface at time \( t \)

\( AB2(t) \) = Concentration of complex (analyte bound to two ligand molecules) at the interaction surface at time \( t \)

\( k_a \) = Association rate constant (single analyte-ligand binding (M\(^{-1}\)s\(^{-1}\)))

\( k_d \) = Dissociation rate constant (single analyte-ligand binding (s\(^{-1}\)))

\( k_{a2} \) = Association rate constant (analyte-ligand complex binding to second ligand molecule (RU\(^{-1}\)s\(^{-1}\)))

\( k_{d2} \) = Dissociation rate constant (analyte-ligand complex binding to second ligand molecule (s\(^{-1}\)))
Two-State Model

This model describes the 1:1 interaction between an analyte and ligand, where the analyte-ligand complex subsequently undergoes a conformational change. Assumptions of this model are that the conformational change only occurs in the complex AB and not in the ligand alone; and that conformationally changed complex does not dissociate freely, but must undergo the reverse conformational change first. The binding model is governed by the following set of differential equations:

\[
\frac{dA}{dt} = -(k_a * A(t) * B - k_d * AB(t)) \\
B(0) = R_{max}
\]

\[
\frac{dAB}{dt} = (k_a * A(t) * B - k_d * AB(t)) - (k_{a2} * AB(t) * B - k_{d2} * AB2(t)) \\
AB(0) = 0
\]

\[
\frac{dAB2}{dt} = k_{a2} * AB(t) - k_{d2} * AB2(t) \\
AB2(0) = 0
\]

where:

- \(A(t)\) = Molar concentration of analyte at the interaction surface at time \(t\)
- \(B\) = Available immobilized ligand
- \(AB(t)\) = Concentration of complex (analyte-ligand binding) at the interaction surface at time \(t\)
- \(AB2(t)\) = Concentration of complex (analyte-ligand in second state) at the interaction surface at time \(t\)
- \(K_a\) = Association rate constant (single analyte-ligand binding (M\(^{-1}\)s\(^{-1}\)))
- \(K_d\) = Dissociation rate constant (single analyte-ligand binding (s\(^{-1}\)))
- \(K_{a2}\) = Association rate constant for formation of conformationally changed complex
- \(K_{d2}\) = Dissociation rate constant for formation of conformationally changed complex
Sales and Service Contacts

Germany
Sartorius Lab Instruments GmbH & Co. KG
Otto-Brenner-Strasse 20
37079 Goettingen
Phone +49 551 308 0

USA
Sartorius Corporation
565 Johnson Avenue
Bohemia, NY 11716
Phone +1 631 254 4249
Or +1 650 322 1360

For further information, visit
www.sartorius.com/octet-support