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1 How to cite these pages

When citing one of these pages, reference the name of the page, the date you accessed it and the URL. All the pages in the statistics and curve fitting guides were written by GraphPad's founder, Harvey Motulsky.

Use this example as a guide to cite a single page:


Note that the URL for each page in these guides is shown at the bottom of the page. Don't use the URL you see in the browser toolbar, as that may be generic and not apply just to the page you are looking at.

Use this example as a guide to citing the entire curve fitting guide:


2 PRINCIPLES OF REGRESSION

Many scientists fit curves more often than the use any other statistical technique. Yet few statistical texts really explain the principles of curve fitting. This Guide provides a concise introduction to fitting curves, especially nonlinear regression.

The first step is to be clear on what your goal is:

- If your goal is to fit a model to your data in order to obtain best-fit values of the parameters, and want to learn the principles first, then read this principles section before trying to fit curves.

- If you already understand the principles of nonlinear regression, and want to see how to fit curves with Prism, jump right to the tutorials.
• If your goal is to simply fit a smooth curve in order to interpolate values from the curve, there is no need to learn much theory. Jump right to an explanation of interpolation with Prism.

• If your goal is to create a spline (a curve that goes through every data point) or a lowess curve (shows the general trend with a curve that can be quite jagged), you can jump right to the instructions for that analysis.

2.1 Understanding mathematical models

A mathematical model is an equation that describes a physical, chemical or biological state or process. The goal of nonlinear regression is to fit a model to your data.

• What exactly is a mathematical model?

• Learn from three example models.

• It sure would be nice if models could be chosen by Prism. Why is this not possible?

• When trying to understand a model, here are some tips for figuring out what it means.

• In addition to fitting a model to data, Prism can plot a function, or a family of functions. You choose a function and values for the parameters, and Prism will plot predicted Y values for a range of X values.

2.1.1 What is a model?

The whole point of nonlinear regression is to fit a model to your data. So that raises the question: What is a model?

A mathematical model is a description of a physical, chemical or biological state or process. Using a model can help you think about chemical and physiological processes or mechanisms, so you can design better experiments and comprehend the results. When you fit a model to
your data, you obtain best-fit values that you can interpret in the context of the model.

A mathematical model is neither a hypothesis nor a theory. Unlike scientific hypotheses, a model is not verifiable directly by an experiment. For all models are both true and false.... The validation of a model is not that it is "true" but that it generates good testable hypotheses relevant to important problems.


Your goal in using a model is not necessarily to describe your system perfectly. A perfect model may have too many parameters to be useful. Rather, your goal is to find as simple a model as possible that comes close to describing your system. You want a model to be simple enough so you can fit the model to data, but complicated enough to fit your data well and give you parameters that help you understand the system, reach valid scientific conclusions, and design new experiments.

2.1.2 Three example models

To give you a sense of how mathematical models work, below is a brief description of three commonly used models.

Optical density as a function of concentration

Background

Colorimetric chemical assays are based on a simple principle. Add appropriate reactants to your samples to initiate a chemical reaction whose product is colored. When you terminate the reaction, the concentration of colored product is proportional to the initial concentration of the substance you want to assay.

Model

Since optical density is proportional to the concentration of colored substances, the optical density will also be proportional to the concentration of the substance you are assaying.

\[ \text{Optical Density} = Y = k \cdot \text{[substance]} = K \cdot X \]
Reality check

Mathematically, the equation works for any value of X. However, the results only make sense with certain values.

- Negative X values are meaningless, as concentrations cannot be negative.
- The model may fail at high concentrations of substance where the reaction is no longer limited by the concentration of substance.
- The model may also fail at high concentrations if the solution becomes so dark (the optical density is so high) that little light reaches the detector. At that point, the noise of the instrument may exceed the signal.

It is not unusual for a model to work only for a certain range of values. You just have to be aware of the limitations, and not try to use the model outside of its useful range.

Exponential decay

Exponential equations whenever the rate at which something happens is proportional to the amount which is left. Examples include ligands dissociating from receptors, decay of radioactive isotopes, and metabolism of drugs. Expressed as a differential equation:

\[
\frac{\Delta Y}{\Delta X} = \frac{dY}{dx} = -k \cdot Y
\]
Converting the differential equation into a model that defines $Y$ at various times requires some calculus. There is only one function whose derivative is proportional to $Y$, the exponential function. Integrate both sides of the equation to obtain a new exponential equation that defines $Y$ as a function of $X$ (time), the rate constant $k$, and the value of $Y$ at time zero, $Y_0$.

$$Y = Y_0 \cdot e^{-kX} = Y_0 \cdot \exp(-k \cdot X)$$

**Equilibrium binding**

When a ligand interacts with a receptor, or when a substrate interacts with an enzyme, the binding follows the law of mass action.

$$R + L \overset{k_s}{\underset{k_{fr}}{\rightleftharpoons}} RL$$

You measure the amount of binding, which is the concentration of the RL complex, so plot that on the $Y$ axis. You vary the amount of added ligand, which we can assume is identical to the concentration of free ligand, $L$, so that forms the $X$ axis. Some simple (but tedious) algebra leads to this equation:

$$\text{Specific binding} = Y = \frac{B_{\text{max}} \cdot X}{K_d + X}$$
2.1.3 The problem with choosing models automatically

The goal of nonlinear regression is to fit a model to your data. The program finds the best-fit values of the parameters in the model (perhaps rate constants, affinities, receptor number, etc.) which you can interpret scientifically.

Choosing a model is a scientific decision. You should base your choice on your understanding of chemistry or physiology (or genetics, etc.). The choice should not be based solely on the shape of the graph.

Some programs (not available from GraphPad Software) automatically fit data to thousands of equations and then present you with the equation(s) that fit the data best. Using such a program is appealing because it frees you from the need to choose an equation. The problem is that the program has no understanding of the scientific context of your experiment. The equations that fit the data best are unlikely to correspond to scientifically meaningful models. You will not be able to interpret the best-fit values of the parameters, so the results are unlikely to be useful.

Letting a program choose a model for you can be useful if your goal is to simply create a smooth curve for simulations or interpolations. In these situations, you don't care about the value of the parameters or the meaning of the model. You only care that the curve fit the data well and does not wiggle too much. Avoid this approach when the goal of curve fitting is to fit the data to a model based on chemical, physical, or biological principles. Don't use a computer program as a way to avoid understanding your experimental system, or to avoid making scientific decisions.

2.1.4 Advice: How to understand a model

Encountering an equation causes the brains of many scientists to freeze.
If you are one of these scientists who has trouble thinking about equations, here are some tips to help you understand what an equation means. As an example, let's use the Michaelis-Menten equation that describes enzyme activity as a function of substrate concentration:

\[ Y = \frac{V_{\text{max}} \times X}{(K_m + X)} \]

**Tip 1. Make sure you know the meaning and units of X and Y**

For this example, Y is enzyme activity which can be expressed in various units, depending on the enzyme. X is the substrate concentration in Molar or micromolar or some other unit of concentration.

**Tip 2. Figure out the units of the parameters**

In the example equation, the parameter \( K_m \) is added to \( X \). It only makes sense to add things that are expressed in the same units, so \( K_m \) must be expressed in the same concentration units as \( X \). This means that the units cancel in the term \( X/(K_m + X) \), so \( V_{\text{max}} \) must be expressed in the same units of enzyme activity as \( Y \).

**Tip 3: Figure out the value of Y at extreme values of X**

Since \( X \) is concentration, it cannot be negative. But it can be zero. Substitute \( X=0 \) into the equation, and you will see that \( Y \) is also zero.

Let's also figure out what happens as \( X \) gets very large. As \( X \) gets large compared to \( K_m \), the denominator \( (X+K_m) \) has a value very similar to \( X \). So the ratio \( X/(X+K_m) \) approaches 1.0, and \( Y \) approaches \( V_{\text{max}} \). So the graph of the model must level off at \( Y=V_{\text{max}} \) as \( X \) gets very large.

**Tip 4. Figure out the value of Y at special values of X**

Since \( K_m \) is expressed in the same units as \( X \), you can ask what happens if \( X \) equals \( K_m \)? In that case, the ratio \( X/(K_m + X) \) equals 0.5, so \( Y \) equals half of \( V_{\text{max}} \). This means the \( K_m \) is the concentration of substrate that leads to a velocity equal to half the maximum velocity \( V_{\text{max}} \).
Tip 5. Graph the model with various parameter values

Graphing a family of curves with various values for the parameters can help you visualize what the parameters mean. To do this with Prism, use the analysis "Create a family of theoretical curves".

2.2 Principles of simple linear regression

Linear regression fits a straight line through your data to find the best-fit value of the slope and intercept.

- **What is the goal of linear regression?**
- **How does it work?**
- Linear regression and correlation are often confused. **How are they distinct?**
- Linear regression is just a special case of nonlinear regression. **How do they differ?**
- Linear regression is sometimes used on transformed data to analyze Scatchard, Lineweaver-Burke and similar plots. **Why is this is not a good way to analyze data?**

2.2.1 The goal of linear regression

**What is linear regression?**

Linear regression fits this model to your data:

\[ Y = \text{intercept} + \text{slope} \times X \]
The slope quantifies the steepness of the line. It equals the change in Y for each unit change in X. It is expressed in the units of the Y axis divided by the units of the X axis. If the slope is positive, Y increases as X increases. If the slope is negative, Y decreases as X increases.

The Y intercept is the Y value of the line when X equals zero. It defines the elevation of the line.

Correlation and linear regression are not the same. Review the differences.

**Simple vs. multiple linear regression**

Simple linear regression is shown above. There is only a single X variable. In contrast, multiple linear regression defines Y as a function that includes several X variables.

### 2.2.2 How linear regression works

**How linear regression works. Minimizing sum-of-squares.**

The goal of linear regression is to adjust the values of slope and intercept to find the line that best predicts Y from X. More precisely, the goal of regression is to minimize the sum of the squares of the vertical distances of the points from the line. Why minimize the sum of the squares of the distances? Why not simply minimize the sum of the actual distances?

If the random scatter follows a Gaussian distribution, it is far more likely to have two medium size deviations (say 5 units each) than to have one small deviation (1 unit) and one large (9 units). A procedure that minimized the sum of the absolute value of the distances would have no
preference over a line that was 5 units away from two points and one that was 1 unit away from one point and 9 units from another. The sum of the distances (more precisely, the sum of the absolute value of the distances) is 10 units in each case. A procedure that minimizes the sum of the squares of the distances prefers to be 5 units away from two points (sum-of-squares = 50) rather than 1 unit away from one point and 9 units away from another (sum-of-squares = 82). If the scatter is Gaussian (or nearly so), the line determined by minimizing the sum-of-squares is most likely to be correct.

The calculations are shown in every statistics book, and are entirely standard.

The term "regression"

The term "regression", like many statistical terms, is used in statistics quite differently than it is used in other contexts. The method was first used to examine the relationship between the heights of fathers and sons. The two were related, of course, but the slope is less than 1.0. A tall father tended to have sons shorter than himself; a short father tended to have sons taller than himself. The height of sons regressed to the mean. The term "regression" is now used for many sorts of curve fitting.

The details

The details of linear regression are explained in numerous text books, so will not be detailed here. Everything Prism does is standard. Here is one good source for the basics including confidence and prediction bands.

2.2.3 Comparing linear regression to correlation

Linear regression is distinct from correlation.

What is the goal?

Linear regression finds the best line that predicts Y from X.

Correlation quantifies the degree to which two variables are related. Correlation does not fit a line through the data points. You simply are computing a correlation coefficient (r) that tells you how much one variable tends to change when the other one does. When r is 0.0, there is no relationship. When r is positive, there is a trend that one variable goes up as the other one
goes up. When $r$ is negative, there is a trend that one variable goes up as the other one goes down.

**What kind of data?**

Linear regression is usually used when $X$ is a variable you manipulate (time, concentration, etc.)

Correlation is almost always used when you measure both variables. It rarely is appropriate when one variable is something you experimentally manipulate.

**Does it matter which variable is $X$ and which is $Y$?**

The decision of which variable you call "$X$" and which you call "$Y$" matters in regression, as you'll get a different best-fit line if you swap the two. The line that best predicts $Y$ from $X$ is not the same as the line that predicts $X$ from $Y$ (however both those lines have the same value for $R^2$).

With correlation, you don't have to think about cause and effect. It doesn't matter which of the two variables you call "$X$" and which you call "$Y$". You'll get the same correlation coefficient if you swap the two.

**Assumptions**

With linear regression, the $X$ values can be measured or can be a variable controlled by the experimenter. The $X$ values are not assumed to be sampled from a Gaussian distribution. The distances of the points from the best-fit line is assumed to follow a Gaussian distribution, with the SD of the scatter not related to the $X$ or $Y$ values.

The correlation coefficient itself is simply a way to describe how two variables vary together, so it can be computed and interpreted for any two variables. Further inferences, however, require an additional assumption -- that both $X$ and $Y$ are measured (are interval or ratio variables), and both are sampled from Gaussian distributions. This is called a bivariate Gaussian distribution. If those assumptions are true, then you can interpret the confidence interval of $r$ and the $P$ value testing the null hypothesis that there really is no correlation between the two variables (and any correlation you observed is a consequence of random sampling).

**Relationship between results**

Linear regression quantifies goodness of fit with $r^2$, sometimes shown in uppercase as $R^2$. If you put the same data into correlation (which is rarely appropriate; see above), the square of $r$ from correlation will equal $r^2$ from regression.

Correlation computes the value of the Pearson correlation coefficient, $r$. Its value ranges from -1 to +1.
Comparing linear regression to nonlinear regression

The goal of linear and nonlinear regression

A line is described by a simple equation that calculates Y from X, slope and intercept. The purpose of linear regression is to find values for the slope and intercept that define the line that comes closest to the data.

Nonlinear regression is more general than linear regression and can fit any model (equation) to your data. It finds the values of those parameters that generate the curve that comes closest to the data.

How linear and nonlinear regression work

Both linear and nonlinear regression find the values of the parameters (slope and intercept for linear regression) that make the line or curve come as close as possible to the data. More precisely, the goal is to minimize the sum of the squares of the vertical distances of the points from the line or curve.

Linear regression accomplishes this goal using math that can be completely explained with simple algebra (shown in many statistics books). Put the data in, and the answers come out. There is no chance for ambiguity. You could even do the calculations by hand, if you wanted to.

Nonlinear regression uses a computationally intensive, iterative approach that can only be explained using calculus and matrix algebra. The method requires initial estimated values for each parameter.

Linear regression is a special case of nonlinear regression

Nonlinear regression programs can fit any model, including a linear one. Linear regression is just a special case of nonlinear regression.

Even if your goal is to fit a straight line through your data, there are many situations where it makes sense to choose nonlinear regression rather than linear regression.

Using nonlinear regression to analyze data is only slightly more difficult than using linear regression. Your choice of linear or nonlinear regression should be based on the model you are fitting. Do not use linear regression just to avoid using nonlinear regression. Avoid
transformations such as Scatchard or Lineweaver-Burke transforms whose only goal is to linearize your data.

2.2.5 Advice: Look at the graph

The four graphs below were created by Anscombe (1) to demonstrate the importance of looking at your data. The four data sets, the Anscome's quartet, all have 11 points. You can see at a glance that the data are very different. Surprisingly, the analyses of these four data sets are identical in many ways. When analyzed by linear regression, all these values are identical for all four graphs:

- The mean values of X and Y
- The slopes and intercepts
- \( r^2 \)
- The standard errors and confidence intervals of the slope and intercept
- The standard deviation of the residuals (sy.x)
- The F ratio and P value when testing the null hypothesis that the slope is really 0.0 (horizontal).
- The covariance between slope and intercept (computed by Prism’s nonlinear regression analysis)
Differences seen in linear regression are:

- The confidence bands, shown in yellow below, are identical for three of the four graphs, but different for the fourth (which has different X values).
- The results of a runs test for linearity.
- Normality tests of residuals.


### 2.2.6 Advice: Avoid Scatchard, Lineweaver-Burke and similar transforms

Before analyzing your data with linear regression, stop and ask yourself whether it might make more sense to fit your data with nonlinear regression. If you have transformed nonlinear data to create a linear relationship, you will almost certainly be better off fitting your original data using nonlinear regression.
Before nonlinear regression was readily available, the best way to analyze nonlinear data was to transform the data to create a linear graph, and then analyze the transformed data with linear regression. Examples include Lineweaver-Burke plots of enzyme kinetic data, Scatchard plots of binding data, and logarithmic plots of kinetic data.

These methods are outdated, and should not be used to analyze data

The problem with these methods is that the transformation distorts the experimental error. Linear regression assumes that the scatter of points around the line follows a Gaussian distribution and that the standard deviation is the same at every value of \( X \). These assumptions are rarely true after transforming data. Furthermore, some transformations alter the relationship between \( X \) and \( Y \). For example, in a Scatchard plot the value of \( X \) (bound) is used to calculate \( Y \) (bound/free), and this violates the assumption of linear regression that all uncertainty is in \( Y \) while \( X \) is known precisely. It doesn't make sense to minimize the sum of squares of the vertical distances of points from the line, if the same experimental error appears in both \( X \) and \( Y \) directions.

Since the assumptions of linear regression are violated, the values derived from the slope and intercept of the regression line are not the most accurate determinations of the variables in the model. Considering all the time and effort you put into collecting data, you want to use the best possible technique for analyzing your data. Nonlinear regression produces the most accurate results.

The figure below shows the problem of transforming data. The left panel shows data that follows a rectangular hyperbola (binding isotherm). The right panel is a Scatchard plot of the same data. The solid curve on the left was determined by nonlinear regression. The solid line on the right shows how that same curve would look after a Scatchard transformation. The dotted line shows the linear regression fit of the transformed data. Scatchard plots can be used to determine the receptor number (\( B_{\text{max}} \), determined as the \( X \)-intercept of the linear regression line) and dissociation constant (\( K_d \), determined as the negative reciprocal of the slope). Since the Scatchard transformation amplified and distorted the scatter, the linear regression fit does not yield the most accurate values for \( B_{\text{max}} \) and \( K_d \).
Don't use linear regression just to avoid using nonlinear regression. Fitting curves with nonlinear regression is not difficult.

Although it is usually inappropriate to analyze transformed data, it is often helpful to display data after a linear transformation. Many people find it easier to visually interpret transformed data. This makes sense because the human eye and brain evolved to detect edges (lines) — not to detect rectangular hyperbolas or exponential decay curves. Even if you analyze your data with nonlinear regression, it may make sense to display the results of a linear transformation.

2.3 Getting started with nonlinear regression

Nonlinear regression finds the values of parameters in a nonlinear model that bring the curve defined by that model close to your data points.

- Distinguishing nonlinear regression from other kinds of regression
- The goal of nonlinear regression
- The six steps of nonlinear regression
- Preparing data for nonlinear regression
- Don't fit a model to smoothed data

2.3.1 Distinguishing nonlinear regression from other kinds of regression

Before choosing nonlinear regression, make sure you don't really need another kind of regression. Also read about how nonlinear regression differs from linear regression.
Polynomial regression

A polynomial model has this form: \( Y = A + BX + CX^2 + DX^3 \) ....

Like linear regression, it is possible to fit polynomial models without fussing with initial values. For this reason, some programs (i.e. Excel) can perform polynomial regression, but not nonlinear regression. And some programs have separate modules for fitting data with polynomial and nonlinear regression. Prism fits polynomial models using the same analysis it uses to fit nonlinear models. Polynomial equations are available within Prism's nonlinear regression analysis.

Multiple regression

A multiple regression model has more than one independent (X) variable. Like linear and nonlinear regression, the dependent (Y) variable is a measurement.

GraphPad Prism 6 does not perform multiple regression. But by using column constants, you can effectively fit models with two independent variables in some circumstances.

Logistic regression

A logistic regression model is used when the outcome, the dependent (Y) variable, has only two possible values. Did the person get the disease or not? Did the student graduate or not? There can be one or several independent variables. These independent variables can be a variable like age or blood pressure, or have discrete values to encode which treatment each subject received.

GraphPad Prism 6 does not perform logistic regression.

Proportional hazards regression

A proportional hazards model is used when the outcome is whether or not a one-time event (often death) occurred. One of the independent variables is time, and other independent variables can be used to account for treatment or other variables.

GraphPad Prism does not perform proportional hazards regression.
2.3.2 The goal of nonlinear regression

Nonlinear regression is used for two purposes

Scientists use nonlinear regression with one of two distinct goals:

- To fit a model to your data in order to obtain best-fit values of the parameters, or to compare the fits of alternative models. If this is your goal, you must pick a model (or two alternative models) carefully, and pay attention all the results.

- To simply fit a smooth curve in order to interpolate values from the curve, or perhaps to draw a graph with a smooth curve. If this is your goal, you can assess it purely by looking at the graph of data and curve. There is no need to learn much theory. Jump right to an explanation of interpolation with Prism.

The general idea of regression

Linear regression fits a straight-line model to your data. Nonlinear regression extends this idea to fit any model to your data. Distinguish nonlinear regression from linear regression and from other types of regression.

The goal of linear and nonlinear regression is to adjust the values of the model's parameters to find the line or curve that comes closest to your data. So with linear regression, the goal is to find the best-fit values of the slope and intercept that makes the line come close to the data. With nonlinear regression of a normalized dose-response curve, the goal is to adjust the values of the EC50 (the concentration that provokes a response halfway between the minimum and maximum responses) and the slope of the curve.

More precisely, the goal of regression is to find the values of the parameters that are most likely to be correct. To do this requires making an assumption about the scatter of data around the curve.

Least-squares regression

The most common assumption is that data points are randomly scattered around an ideal curve (or line) with the scatter following a Gaussian distribution. If you accept this assumption, then the goal of regression is to adjust the model's parameters to find the curve that minimizes the sum of the squares of the vertical distances of the points from the curve.
Why minimize the sum of the squares of the distances? Why not simply minimize the sum of the actual distances?

If the random scatter follows a Gaussian distribution, it is far more likely to have two medium size deviations (say 5 units each) than to have one small deviation (1 unit) and one large (9 units). A procedure that minimized the sum of the absolute value of the distances would have no preference over a curve that was 5 units away from two points and one that was 1 unit away from one point and 9 units from another. The sum of the distances (more precisely, the sum of the absolute value of the distances) is 10 units in each case. A procedure that minimizes the sum of the squares of the distances prefers to be 5 units away from two points (sum-of-squares = 25) rather than 1 unit away from one point and 9 units away from another (sum-of-squares = 82). If the scatter is Gaussian (or nearly so), the curve determined by minimizing the sum-of-squares is most likely to be correct.

2.3.3 The six steps of nonlinear regression

Before learning how to fit a model using GraphPad Prism (or any program), it is worth first reviewing the necessary steps.

Step 1. Choose a model

Nonlinear regression fits a model to your data. You must, therefore, choose a model or enter a new model. This is a scientific decision that must be made by someone who understands the scientific goals of the experiment. Why a computer program should not pick a model for you.

Step 2. Decide whether to constrain any parameters to constant values

When performing nonlinear regression, you don't have to fit each parameter in the equation. Instead, you may fix one or more of the parameters to constant values. It is often helpful to define constants when you have only a few data points. For example, you might fix the bottom plateau of a sigmoid curve or exponential decay to zero.

Don't expect computer programs to have common sense. That is your job! Think about how you did the experiment and prepared the data, and decide whether some of the parameters should be fixed. For example, if a background signal has already been subtracted, it makes sense to fix
the bottom plateau of a dose-response curve or an exponential decay curve to zero.

**Step 3. Choose (or review) initial values**

Nonlinear regression is an iterative procedure. The program must start with estimated values for each parameter. It then adjusts these initial values to improve the fit. GraphPad Prism supplies initial values automatically if you choose a built-in equation. If you enter your own equation, you will also need to provide initial values, or rules to generate initial values from the range of the data.

You'll find it easy to estimate initial values if you have looked at a graph of the data, understand the model, and understand the meaning of all the parameters in the equation. Remember that you just need an estimate. It doesn't have to be very accurate. If you are having problems estimating initial values, set aside your data and simulate curves using the model. Change the variables one at a time, and see how they influence the shape of the curve. Once you have a better feel for how the parameters influence the curve, you might find it easier to estimate initial values.

When fitting a simple model to clean data, it won't matter much if the initial values are fairly far from the correct values. You'll get the same best-fit curve no matter what initial values you use, unless the initial values are extremely far from correct. Initial values matter more when your data have a lot of scatter or your model has many variables.

**Step 4. If you are fitting two or more data sets at once, decide whether to share any parameters**

If you enter data into two or more data set columns, Prism will fit them all in one analysis. But each fit will be independent of the others unless you specify that one or more parameters are shared. When you share parameters, the analysis is called a global nonlinear regression.

**Step 5. Decide on a weighting scheme**

Nonlinear regression programs generally weight each point equally. But there are many ways to differentially weight the points.
Step 6. Choose which results you want reported

Nonlinear regression always reports the best-fit values of the parameters. Beyond that, Prism (and most programs) gives you many choices for which results you want reported.

2.3.4 Preparing data for nonlinear regression

You must create an XY data table in Prism, for use with nonlinear regression.

Follow these guidelines to enter (or preprocess) data for nonlinear regression:

- **Avoid linearizing transforms** such as Scatchard and Lineweaver-Burke plots. Such plots are useful for displaying data but are obsolete for data analysis.

- Transforming X values can be convenient, and will not change the results of regression (so long as the model is adjusted accordingly). Use Prism's Transform analysis to do this.

- **Don't smooth your data**. You will get invalid nonlinear regression results. Fit the raw data.

- Transforming Y values to change units or to subtract a baseline can be convenient, and will not substantially affect nonlinear regression. Use Prism's Transform analysis to do this.

- Avoid nonlinear Y transforms (reciprocals, logs) unless you have a very good reason. Such a transform can be useful if it equalizes the variances (so the scatter at all points along the curve is about the same), but should not be done just to linearize the data.

- Enter raw replicates when possible, and not just mean and SD or SEM. Prism will fit the same curve either way, but there are two advantages of entering the raw data. First, you can plot every individual replicate so you see the actual data. Second, entering the
individual replicates lets you choose robust nonlinear regression or automatic outlier elimination

- If you have entered replicates, first graph the individual replicates rather than mean and error bar. You may want to later plot mean and error bar, but look at a graph of the raw data first.

2.3.5 Don’t fit a model to smoothed data

The figure shows the number of hurricanes over time. The left panel shows the number of hurricanes in each year, which jumps around a lot. To make it easier to spot trends, the right panel shows a rolling average. The value plotted for each year is the average of the number of hurricanes for that year plus the prior eight years. This smoothing lets you see a clear trend.

But there is a problem. These are not real data. Instead, the values plotted in the left panel were chosen randomly (from a Poisson distribution, with a mean of 10). There is no pattern. Each value was randomly generated without regard to the previous (or later) values.

Creating the running average creates the impression of trends by ensuring that any large random swing to a high or low value is amplified, while the year-to-year variability is muted.

You should not fit a model to the rolling average data with linear or nonlinear regression, or compute a correlation coefficient. Any such results would be invalid and misleading. The problem is that regression assumes that each value is independent of the others, but the
rolling average are not at all independent of each other. Rather each value is included as part of the neighboring values.

This example is adapted from Briggs (2008).

2.3.6 Reparameterizing an equation can help

Even though nonlinear regression, as its name implies, is designed to fit nonlinear models, some of the inferences actually assume that some aspects of the model are close to linear, so that the uncertainty about each parameter's value is symmetrical.

Reparameterizing an equation can make the uncertainty more symmetrical, making the SE easier to interpret and making the symmetrical asymptotic CI more helpful. Prism 7 now can compute an asymmetrical CI, and when you choose this approach it doesn't matter so much how you parametrize the equation.

What is reparameterizing?

There are two forms of the model used to fit the sigmoidal enzyme kinetics data to a standard model:

\[ Y = \frac{V_{max} \times X^h}{K_{half}^h + X^h} \]
\[ Y = \frac{V_{max} \times X^h}{K_{prime} + X^h} \]

The two equations are equivalent. They both fit Vmax (the maximum activity extrapolated to very high concentrations of substrate) and h (Hill slope, describing the steepness of the curve). But one model fits Khalf (the concentration needed to obtain a velocity half of maximal) and the other fits Kprime (a more abstract measure of substrate action).

Which model is best? The two are equivalent, with Kprime equal to Khalf^h, so they will generate exactly the same curve.
Since the sum-of-squares will be identical, as will the number of degrees of freedom, any comparison of this model with another will give identical results regardless of which form of this model you pick.

**Distribution of parameters are not always symmetrical**

Simulations can determine how symmetrical a parameter's uncertainty is. I simulated sigmoidal enzyme kinetics using $V_{\text{max}}=100$, $h=5$, $K_{\text{prime}}=25$ (so $K_{\text{half}}=5$), and Gaussian scatter with a SD equal to 7.5. The X values matched those in the figure above, with triplicate Y values at each X. Prism can repeat such simulations easily. I repeated the simulations 5000 times, fit each curve to both forms of the models, and tabulated the best-fit values of $K_{\text{prime}}$ and $K_{\text{half}}$, and computed the skewness of each.
Clearly the distribution of Khalf is quite symmetrical, and looks Gaussian. The skewness is close to zero, as expected for a symmetrical distribution. In contrast, the distribution of Kprime is quite skewed. Note that a few of the simulations had best-fit values of Kprime greater than 100. The skewness value (4.89) confirms what is obvious by inspection -- the distribution is far from symmetrical.

Consequence of asymmetrical parameters

Ideally, a confidence interval is easy to interpret. A 95% CI has a 95% chance of including the true population value of the parameter, and a 5% chance of missing it.

When analyzing real data, we never know the value of the true parameter, so never know if the interval includes it or not. But when you simulate data, you know the true values of the parameters, so can quantify the coverage of the confidence intervals. I set up the same simulations mentioned above, fit each data set to both equations, and tabulated whether each confidence interval included the true parameter value or not. This table shows the fraction of 5,000 simulations where the asymptotic symmetrical confidence interval did not include the true parameter value (25 for Kprime, and 1.9037 for Khalf).

<table>
<thead>
<tr>
<th></th>
<th>&quot;95% CI&quot;</th>
<th>&quot;99% CI&quot;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ideal</td>
<td>5.0%</td>
<td>1.0%</td>
</tr>
</tbody>
</table>
These results show that Khalf is well behaved, as expected given its symmetry (see above). The 95% confidence interval is expected to miss the true value in 5.0% of the simulations. In fact, it happened 5.1% of the time. Similarly, the 99% CI is expected to miss the true value in 1.0% of the simulations, which is exactly what happened. In contrast, Kprime is less well behaved. The intervals computed to be 95% confidence intervals were not wide enough so missed the true value in 8.8% of the simulations. The 99% intervals were similarly not wide enough so missed the true value in 4.8% of the simulations. Thus the confidence intervals computed to be 99% intervals, actually turned out to be 95% intervals.

These simulations show the advantage of choosing the equation that fits Khalf, rather than the one that fits Kprime. Khalf has a symmetrical distribution so the confidence intervals computed from these fits can be interpreted at face value. In contrast, Kprime has an asymmetrical distribution and its confidence intervals cannot be interpreted at face value.

**Parameterization doesn't matter if you ask Prism to present the asymmetrical likelihood ratio confidence intervals**

If you choose asymmetrical likelihood ratio confidence intervals (new in Prism 7), then it doesn't matter which form of the equation you choose. The coverage will be the same in both cases, and will be very close to 95% or 99%. With this choice, you can choose the form of the equation that matches text books and papers, or that fits the way you think. If you prefer to think graphically, choose the Khalf. If you think mechanistically, choose Kprime.

2.4 **Weighted nonlinear regression**
2.4.1 The need for unequal weighting in nonlinear regression

Both graphs above show dose-response curves, with response measured in ten replicate values at each response. In the left graph, the standard deviation of those replicates is consistent. It is about the same all the way along the curve. In the right graph, the standard deviation of the replicates is related to the value of Y. As the curve goes up, variation among replicates increases.

These data are simulated. In both cases, the scatter among replicates is sampled from a Gaussian distribution. In the graph on the left, the SD of that Gaussian distribution is the same for all doses. On the right, the SD is a constant fraction of the mean Y value. When a response is twice as high as another response, the standard deviation among replicates is also twice as large. In other words, the coefficient of variation (CV) is constant.

What happens if you fit a model to the data on the right without taking into account the fact that the scatter increases as Y increases? Consider two doses that give different responses that differ by a factor of two. The average distance of the replicates from the true curve will be twice as large for the higher response. Since regression minimizes the sum of the squares of those distances those points will be expected to contribute four times as much to the sum-of-squares as the points with the smaller average Y value. In other words, the set of replicates whose average Y value is twice that of another set will be given four times as much weight. This means, essentially, that the curve fitting procedure will work harder to bring the curve near these points, and relatively ignore the points with lower Y values. You’d need to have four times as many replicates in the lower set to equalize the contribution to the sum-of-squares.
The goal of weighting is for points anywhere on the curve to contribute equally to the sum-of-squares. Of course random factors will give some points more scatter than others. But the goal of weighting is to make those differences be entirely random, not related to the value of Y. The term weight is a bit misleading, since the goal is to remove that extra weight from the points with high Y values. The goal is really to unweight.

Prism offers six choices on the Weights tab of nonlinear regression, and lets you test for appropriate weighting.

2.4.2 Math theory of weighting

Relative weighting (weighting by 1/Y2)

In many experimental situations you expect the average distance (or rather the average absolute value of the distance) of the points from the curve to be higher when Y is higher. Another way to say this is that you expect the standard deviation of the residuals (distance of points from the curve) to get larger when Y is higher -- for the standard deviation to be proportional to the value of Y. Yet another way to say this is that you expect the coefficient of variation to be constant, but for the SD to not be constant.

If the SD of residuals is related to the value of Y, then the points with larger Y values will tend to be further from the curve (have larger residuals). These points will therefore contribute more to the sum-of-squares, and thus dominate the calculations.

The solution is not to minimize the sum-of-squares, but rather to minimize the weighted sum-of-squares. In other words, minimize the relative distances of the Y values of the data (Ydata) from the Y values of the curve (Ycurve). When you choose relative weighting, nonlinear regression minimizes this quantity:

$$\sum \left( \frac{Y_{data} - Y_{curve}}{Y_{curve}} \right)^2 = \sum \left( \frac{Y_{data} - Y_{curve}}{Y_{curve}} \right)^2$$

The left side is easiest to understand. For each point, compute how far it is (in the Y direction) from the curve, divide that value by the Y value of
the curve, and square that ratio. Add up that squared ratio for all points. The right side is equivalent. It squares the numerator and denominator separately, and then computes the ratio of those two squared values. This is the way most mathematical statisticians think about weighting, so relative weighting is also called weighting by $Y^2$.

**Poisson weighting**

Weighting by $1/Y$ is a compromise between minimizing the actual distance squared and minimizing the relative distance squared. One situation where $1/Y$ weighting is appropriate is when the Y values follow a Poisson distribution. This would be the case when Y values are radioactive counts and most of the scatter is due to counting error. With the Poisson distribution, the standard error of a value equals the square root of that value. Therefore you divide the distance between the data and the curve by the square root of the value, and then square that result. The equation below shows the quantity that Prism minimizes, and shows why it is called weighting by $1/Y$ (but Prism actually weights by the absolute value of Y.)

$$\sum \left( \frac{Y_{data} - Y_{curve}}{\sqrt{Y_{curve}}} \right)^2 = \sum \left( \frac{Y_{data} - Y_{curve}}{|Y_{curve}|} \right)^2$$

**General weighting**

The first three equations below shows how absolute, Poisson and relative weighting are related. Note that taking anything to the zero power results in 1.0, so the denominator of equation on the left always equals 1.0.

<table>
<thead>
<tr>
<th>Absolute</th>
<th>Poisson</th>
<th>Relative</th>
<th>General</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\sum \left( \frac{Y_{data} - Y_{curve}}{Y_{curve}} \right)^2$</td>
<td>$\sum \left( \frac{Y_{data} - Y_{curve}}{Y_{curve}} \right)^2$</td>
<td>$\sum \left( \frac{Y_{data} - Y_{curve}}{Y_{curve}} \right)^2$</td>
<td>$\sum \left( \frac{Y_{data} - Y_{curve}}{Y_{curve}} \right)^2$</td>
</tr>
<tr>
<td>$K=0$</td>
<td>$K=1$</td>
<td>$K=2$</td>
<td>$K=???$ (you enter)</td>
</tr>
</tbody>
</table>
The equation on the right shows general weighting. You enter K so can customize the weighting scheme to fit your data. Generally this choice is used with values of K between 1.0 and 2.0. Reference 1 below uses this approach.

If you want to experimentally determine the best value of K, you can do so:

1. Collect data with lots (over a dozen; maybe several dozen) replicates at many points along the curve.

2. Plot the data the usual way to make sure the data seem correct.

3. Create a second graph that ignores the X values (time or concentration...). Instead, in this new graph, X is the logarithm of mean of the replicate Y values for each point on the first graph, and Y on this new graph is the logarithm of variance (square of the standard deviation) among the replicate values. You can use either natural logarithms or logarithms base 10. It doesn't matter, so long as both logarithms uses the same base.

4. Fit a straight line to this graph using linear regression. Since the assumption of a Gaussian variation around this line is dubious, use nonlinear regression and choose a robust fit.

5. The slope of this regression line is K. If K is close to 0.0, then the SD does not vary with Y so no weighting is needed. If K is close to 2.0, then the standard deviation is proportional to Y, and relative weighting is appropriate. If K is close to 1.0, choose Poisson weighting. If K has some other value, use general weighting and enter this value of K as a constant.

Note that if Ycurve is negative, Prism actually takes the the absolute value of Ycurve to the K power.

**Rarely used weighting choices in Prism**

The choices to weight by 1/X or 1/X² weight the points at the left part of the graph more than points to the right. This is common in some fields when fitting straight lines to bioassay data.

Prism also offers the choice to weight by the reciprocal of the standard deviation squared. This option minimizes the following:
This method is most useful when you have formatted the data table for entry of SD values, but then entered values that you had calculated elsewhere based on understanding how the scatter (or errors) arise in your experimental system. The "SD" values you enter should be computed weighting factors, not the actual SD of the data.

If you enter the actual SD into the SD subcolumn, or enter replicate values so Prism computes the SD, then Prism will use these actual SD values as weighting factors. This is way less useful than it sounds. With small to moderate sample sizes, the SD will jump around a lot by chance, and it is not appropriate to use these random SD values for weighting. Weighting should be by predicted SD not actual SD (which is subject to random factors).

Of course, weighting by SD is impossible if any of the entered SD values are zero. It also is impossible if Prism is computing the SD from replicates and there is only one replicate, or if all replicates are identical (so their SD equals zero).

**Prism doesn't use weighting on the first iteration**

There is one situation where weighting by the predicted Y values of the curve causes problems - when the initial values are really bad. The initial curve can be quite far from the points, and in some cases can have Y=0 for some X values, which makes weighting by 1/Y or 1/Y² impossible. To improve the fitting, even if the initial values generate a curve far from the data, Prism uses no weights for the first iteration. This first iteration brings the curve closer to the points. From then on, Prism uses the weighting function you specify. Essentially it uses the results of one iteration of unweighted fitting as the initial values for the weighted fit.

**Reference**

2.4.3 Don't use weighted regression with normalized data

Before fitting a log(dose) vs. response curve, it is common to first normalize the data. This page explains why one should not use weighted regression in this situation.

Weighted fits to nonnormalized data -- works great

The graph below shows simulated data that show a common situation. The scatter among replicates is greater when the Y values are greater. In fact, these data were simulated so the SD among replicates is proportional to the Y value.

These data fit nicely using relative weighting. This minimizes the sum of the squares of the relative distance between the points and the curve. In other words it minimizes:

\[ \sum \left( \frac{Y_{\text{data}} - Y_{\text{curve}}}{Y_{\text{curve}}} \right)^2 = \sum \left( \frac{Y_{\text{data}} - Y_{\text{curve}}}{Y_{\text{curve}}} \right)^2 \]
I ran 10,000 simulations, and found that in every case the fit worked well and gave a reasonable answer (with the EC50 within the range of the data). No surprise so far. The data were fitting assuming a model that exactly matched the method used to simulate the data, and those fits worked well.

**Weighted fits to normalized data -- ugh**

People often like to normalize dose-response data so the Y values range from 0% to 100%. What happens if these normalized data are fit with weighted nonlinear regression?

I answered that question with simulations. Of 1,000 simulated data sets, 223 could not be fit at all. Moreover, 60 simulated data sets gave nonsense results, with the EC50 outside of the range of the data. The remaining 72% of the simulations seemed ok, but the confidence intervals were very wide in some.

**What's going on?**

With the data that were not normalized, the Y value of the best-fit curve at X = -4 is 3.05 times larger than the Y value of the curve at X=-8. Since the weighting is proportional the the Y value of the curve squared, the points near the top of the curve (at X=-4) get $3.05^2$, or 9.28, times as much weight as points near the bottom of the curve (at X=-8).

The situation is very different with the normalized data. With the first simulated data set, the Y value of the best-fit curve at X = -4 is 17.77 times larger than the Y value of the curve at X=-8. Since the weighting is
proportional the Y value of the curve squared, the points near the top of the curve (at X=-4) get $17.77^2$, or 315.8, times as much weight as points near the bottom of the curve (at X=-8). Since the points at the top of the curve gets so much more weight that those at the bottom, those at the bottom are essentially ignored, making the entire curve fit not work very well.

Another problem. Some of the Y values near the bottom of the normalized data sets are negative. Weighting factors really don't make any sense at all when some values are negative and some are positive.

In summary: After normalizing the SD among replicates is no longer proportional to the Y values, so relative weighting is not appropriate.

**Bottom line**

The whole idea of weighted nonlinear regression is to match the weighting scheme used by the regression to the variation in the actual data. If you normalize the data, none of the usual weighting schemes will work well.

If you really want to show your data on a normalized axis running from 0% to 100%, you can do so. First fit the model to the actual data using an appropriate weighting scheme. Then normalize both the data and the curve.

[Details and link to Prism file](#)

### 2.4.4 What are the consequences of choosing the wrong weighting method

What happens if you leave the choice on the weighting tab set to its default -- no weighting -- when the scatter is actually related to the Y value? Of course, the answer is "it depends"? It depends on how large the scatter is, and how large a range the Y values span. The rest of this article looks at one example, and simulates many data sets to see what happens when the fitting is done inappropriately.

**Simulations**

I picked a very simple model -- a straight line. I simulated the data so the SD of scatter at any point along the line is proportional to the Y value
of that point. The graph below shows a simulated data set. You can clearly see that the scatter among replicates increases as the line goes up.

The line was fit to the data by "nonlinear" regression. Prism does not offer differential weighting as part of its linear regression analysis, but "nonlinear" regression can fit a straight line with many options not available in the linear regression analysis.

The red line used the default choice -- no weighting; minimize sum of squares. The blue line used relative weighting. This choice is appropriate when you expect the SD of replicate residuals to be proportional to Y. The two lines are not identical.

I simulated 5000 such data sets using the Monte Carlo analysis of Prism 6. Each of the 5000 simulated data sets was fit with both unweighted and weighted (relative weighting) regression. I recorded both the best-fit value of the slope and its standard error (SE) for both analyses of each of the 5000 data sets.
The scatter among replicates was proportional to Y, so relative weighting is appropriate. The results of these simulations are shown in blue. The results in red show the results when the regression used equal weighting (the default). The boxes extend from the 25th to the 75th percentile, with a line at the median (50th percentile). The whiskers extend down to the smallest value and up to the largest.

The left graph above shows the distribution of the best-fit values of the slopes. Choosing the wrong weighting scheme would not systematically result in a best-fit value of slope that is too high (too steep) or too low (too shallow). Indeed, the median best-fit values for slope are almost identical for the no-weighting and relative-weighting fits. But note that the results with no weighting has a much broader spread of slopes. Since these are simulated data, we know the true slope (1.0) and can quantify how far the best-fit slope is from that idea for each simulated data set. The median error when the data were fit without differential weighting (red) was 0.04610, compared to 0.02689 for the simulations with relative weighting (blue). In this example, choosing the wrong weighting scheme increased the median error by 71%.

The graph on the right above shows the distribution of the SE of the slopes. The SE of the slope is much higher on average and less consistent (wider spread) when the fits were done with equal weighting for all points. The median SE for the unweighted fits was 73% larger than the median SE for the weighted fits. Since the width of the confidence
interval is proportional to the SE, this means the median confidence interval was 73% wider when the fits were done without weighting.

Since these are simulated data, we know the true population slope (it is 1.0). So we can ask for each simulation whether the reported 95% confidence interval included the true value. For the relative weighting simulations, the 95% confidence intervals included the true value in 95.3% of the simulated data sets. When those same data sets were analyzed without weighting, only 92.6% of the "95%" confidence intervals included the true value.

**Summary**

In summary, when we chose the wrong weighting scheme in this example:

- The best-fit value of the slope was less precise.

- The SE of the slope was larger, and thus the confidence intervals were wider.

- Even though the 95% confidence intervals were wider, they weren't wide enough. The "95%" confidence interval only included the true value in fewer than 95% of the simulations.

This is just one example. With other examples, the choice of weighting method matters less. But with still other examples, it may matter more. It is worth trying to choose the appropriate weighting scheme.

[Download the Prism file for this example.](#)

### 2.4.5 Poisson regression

### 2.5 The many uses of global nonlinear regression

A global model defines a family of curves, rather than just a single curve, with some parameters shared between data sets. For each shared parameter, Prism fits one (global) best-fit value that applies to all the data sets. For each non-shared parameter, Prism fits a separate (local) best-fit value for each data set.
2.5.1 What is global nonlinear regression?

A global model defines a family of curves, rather than just a single curve. Some parameters are shared so a single parameter value applies to all the curves, while other parameters apply to each data set individually. For each shared parameter, fit one (global) best-fit value that applies to all the data sets. For each non-shared parameter, fit a separate (local) best-fit value for each data set.

Nonlinear regression finds parameters of a model that make the curve come as close as possible to the data. This is done by minimizing the sum of the squares of the vertical distances between the data points and curve. Global nonlinear regression extends this idea to fitting several data sets at once and minimizes the sum (of all data sets) of sum (of all data points) of squares.

Prism makes it very easy to perform global nonlinear regression. Enter your data on one data table, click analyze, choose nonlinear regression and choose a model. On the Constrain tab of the Nonlinear regression dialog, choose which parameter(s) to share among data sets.

2.5.2 The uses of global nonlinear regression

Global nonlinear regression is useful in (at least) three situations:

- Test whether a parameter value differs significantly between data sets. Compare the sum-of-squares (to assess goodness-of-fit) when the parameter is shared, with the sum of the sum-of-squares when the parameter is fit individually to each dataset. In Prism, set up this kind of comparison in the Compare tab.
• Fit families of data where each dataset is incomplete, but the entire family of datasets defines the parameters. For example, one data set may do a great job of defining the bottom plateau of a dose-response curve, while another data set defines the top. Fit the two data sets separately, and the results may be ambiguous (very wide confidence intervals). Globally fit both curves, and the results might be very tight. See an example.

• Fit models where the parameter(s) you care about cannot be determined from any one dataset, but only from the relationship between several data sets. Another example is fitting enzyme inhibition data.

The first two uses of global fitting do not require writing special models. The third use requires that you write a model for this purpose.

2.5.3 Using global regression to fit incomplete datasets

The graph below shows two dose-response curves. The goal of the experiment is to determine the two EC$_{50}$ values. The EC$_{50}$ is the concentration (dose) that gives a response half-way between the minimum and maximum responses. Each curve in the graph below was fit individually to one of the data sets. The horizontal lines show the 95% confidence interval of the EC$_{50}$.

While the curves nicely fit the data points, the confidence intervals are quite wide. We really haven't determined the EC$_{50}$ with sufficient precision to make useful conclusions. The problem is that the control data (squares) don’t really define the bottom plateau of the curve, and the treated data (circles) don’t really define the top plateau of the curve. Since the data don’t define the minimum and maximum responses very well, the data also don’t define very clearly the point half-way between
the minimum and maximum responses. Accordingly, the confidence intervals for each EC\textsubscript{50} extend over more than an order of magnitude. The whole point of the experiment was to determine the two EC\textsubscript{50} values, but there is an unacceptable amount of uncertainty in the value of the best-fit values of the EC\textsubscript{50}.

The problem is solved by sharing parameters. For this example, share the parameters that define the top and bottom plateaus and the slope. But don’t share the EC50 value, since the EC50 values for control and treated data are clearly distinct.

Here are the results.

The graph of the curves looks only slightly different. But now the program finds the best-fit parameters with great confidence. The 95% confidence intervals for the EC\textsubscript{50} values span about a factor of two (compared to a factor of ten or more when no parameters were shared).

The control data define the top of the curve pretty well, but not the bottom. The treated data define the bottom of the curve pretty well, but not the top. By fitting both data sets at once, sharing some parameters, both EC50 values were determined with reasonable certainty.

2.5.4 Fitting models where the parameters are defined by multiple data sets

Global fitting is most useful when the parameters you care most about are not defined by any one data set, but rather by the relationship between two data sets.
Sample data

Choose the XY sample data set: Binding --Saturation binding to total and nonspecific

Fit the data using nonlinear regression, open the "Binding -- Saturation" list of equations, and choose "One site -- total and nonspecific". You'll see the fit below.

![Graph showing total and nonspecific binding](image)

Explanation of the equation and global fitting

This experiment measured equilibrium binding of radioligand at various concentrations of radioligand to find the $B_{\text{max}}$ and $K_d$ of the radioligand. Since the ligand binds to nonspecific sites as well as the receptor of interest, the experiment measured both total binding and nonspecific binding (binding of radioligand in the presence of an excess of an unlabeled receptor blocker).

These kind of data are often analyzed by first subtracting the nonspecific binding from the total binding. The resulting specific binding is then fit to a model that describes equilibrium binding to one receptor site.

Global fitting simultaneously fits both the total binding and the nonspecific binding. There is no need to first subtract the two data sets. The only trick is to write a model that fits different equations to each data set. Prism's built in equation is set up as follows:
specific=Bmax*X/(X+Kd)
nonspecific=NS*X + Background

\[ Y = \text{specific} + \text{nonspecific} \]
\[ Y = \text{nonspecific} \]

The first line defines specific saturable binding.

The second line defines nonspecific binding to be a constant fraction of added radioligand (X) plus a background (which is often zero).

The third line is preceded by \(<A>\), so it only applies to the first data set (column A, total binding). It defines the Y values in that dataset to equal the sum of total and nonspecific binding.

The fourth line is preceded by \(<B>\) so only applies to the second data set, and defines those Y values to equal nonspecific binding.

The equation is defined with the constraint that the parameters NS and background are shared between the two data sets. That way, Prism finds one best-fit value for NS and background, based on fitting both data sets. Since Bmax and Kd are only used in fitting the first dataset, it wouldn't be meaningful to share these parameters.

The parameters you care about (Bmax and Kd) cannot be determined precisely by fitting just one dataset. But fitting a model that defines both data sets (and their relationship) while sharing the parameter NS between the datasets, lets Prism get the most information possible from the data.

2.5.5 Advice: Don’t use global regression if datasets use different units

Global fitting works by minimizing the sum (for all data sets) of the sum (for all data points) of squares of distances of the data points from the curve. This approach only makes sense when all the data are expressed in the same units.

If different data sets are expressed in different units, be very cautious about using global fitting. The problem is that your decision about which units to use can change the results.

For example, imagine what happens if you change one data set from expressing weight in grams to expressing weight in milligrams. All the
values are now increased by a factor of one thousand, and the sum-of-squares for that data set is increased by a factor of one million (one thousand squared). Compared to other data sets, expressed in different units, this data set now has a much greater impact on the fit. If you really need to do global fit to data sets using different units, consider first normalizing the data so they are comparable.

### 2.6 Comparing fits of nonlinear models

When fitting a nonlinear model to data, your main objective is often to discriminate between different models, or to ask whether an experimental intervention changed a parameter.

- Comparing models can answer **four distinct questions**:
  - For each data set, which of two equations (models) fits best?
  - Do the best-fit values of selected unshared parameters differ between data sets?
  - For each dataset, does the best-fit value of a parameter differ from a theoretical value?
  - Does one curve adequately fit all the data?

- Prism lets you choose from **two approaches to comparing models**: the extra sum-of-squares F tests and the AICc approach based on information theory.

- **How the F test works to compare models**

- **How the AICc computations work**

- The idea of comparing models extends way beyond nonlinear regression. In fact, much of statistics can be viewed as comparing models...
2.6.1 Questions that can be answered by comparing models

When fitting data with regression, in many cases your main objective may be to compare the fits of different models, or to ask if an experimental intervention changed a parameter.

On the Compare tab of the Nonlinear regression dialog, Prism offers four choices:

**For each data set, which of two equations (models) fits best?**

Compare the fit of two models, taking into account differences in the number of parameters to be fit. Most often, you will want to compare two related equations. Comparing the fits of two unrelated equations is rarely helpful.

Example: Compare a one-phase exponential decay with a two-phase exponential decay.

**Do the best-fit values of selected unshared parameters differ between data sets?**

Compare the fit when the selected parameter(s) are shared among all datasets with the fit when those parameter(s) are fit individually to each dataset.

If you pick one parameter, you are asking whether the best-fit value of that one parameter differs among datasets.

If you pick all the parameters, you are asking whether a single curve adequately fits all the data points, or if you get a better fit with individual curves for each dataset.

Example: Fit a family of dose-response curves and compare the fit when the slope factor (Hill slope) is shared with the fit when each curve is fit individually. This is a way to test whether the curves are parallel.

**For each dataset, does the best-fit value of a parameter differ from a theoretical value?**

You may have theoretical reasons to believe that a parameter will have a certain value (often 0.0, 100, or 1.0). Compare the fit when the parameter is constrained to that value with the unconstrained fit.
Example: Test if a Hill Slope differs from 1.0 (a standard value).

Does one curve adequately fit all data sets?

This choice compares the fits of separate curves to each data set with the fit of a single curve fit to all the data sets. It asks whether there is evidence that the treatments did anything to shift the curves.

This choice is new in Prism 6 and is identical to choosing "Do the best-fit values of selected unshared parameters differ between data sets?" and then selecting all the parameters.

2.6.2 Approaches to comparing models

Approach to comparing models

Which model is 'best'? At first, the answer seems simple. The goal of nonlinear regression is to minimize the sum-of-squares, so it seems as though the model with the smaller sum-of-squares is best. If the two alternative models both have the same numbers of parameters, then indeed that is the best approach.

But that approach is too simple when the models have different numbers of parameters, which is usually the case. A model with more parameters can have more inflection points, so of course comes closer to the points. It can bend and twist more to get near the data points. Accordingly, a two-phase model almost always fits better than a one-phase model, and a three-phase model fits even better. So any method to compare a simple model with a more complicated model has to balance the decrease in sum-of-squares with the increase in the number of parameters.

Two statistical approaches to comparing models

Prism offers two approaches to comparing models with different numbers of parameters. These are not the only methods that have been developed to solve this problem, but are the most commonly used methods.

Extra sum-of-squares F test

The Extra sum-of-squares F test is based on traditional statistical hypothesis testing.
The null hypothesis is that the simpler model (the one with fewer parameters) is correct. The improvement of the more complicated model is quantified as the difference in sum-of-squares. You expect some improvement just by chance, and the amount you expect by chance is determined by the number of data points and the number of parameters in each model. The F test compares the difference in sum-of-squares with the difference you would expect by chance. The result is expressed as the F ratio, from which a P value is calculated.

The P value answers this question:

If the null hypothesis is really correct, in what fraction of experiments (the size of yours) will the difference in sum-of-squares be as large as you observed, or even larger?

If the P value is small, conclude that the simple model (the null hypothesis) is wrong, and accept the more complicated model. Usually the threshold P value is set at its traditional value of 0.05. If the P value is less than 0.05, then you reject the simpler (null) model and conclude that the more complicated model fits significantly better.

Information theory approach Akaike's criterion (AIC)

This alternative approach is based on information theory, and does not use the traditional “hypothesis testing” statistical paradigm. Therefore it does not generate a P value, does not reach conclusions about “statistical significance”, and does not “reject” any model.

The method determines how well the data supports each model, taking into account both the goodness-of-fit (sum-of-squares) and the number of parameters in the model. The results are expressed as the probability that each model is correct, with the probabilities summing to 100%. If one model is much more likely to be correct than the other (say, 1% vs. 99%), you will want to choose it. If the difference in likelihood is not very big (say, 40% vs. 60%), you will know that either model might be correct, so will want to collect more data. How the calculations work.

Which approach to choose?

In most cases, the models you want to compare will be 'nested'. This means that one model is a simpler case of the other. For example, a one-phase exponential model is a simpler case of a two-phase exponential model. A three parameter dose-response curve with a standard Hill slope
of 1.0 is a special case of a four parameter dose-response curve that
finds the best-fit value of the Hill slope as well.

If the two models are nested, you may use either the F test or the AIC
approach. The choice is usually a matter of personal preference and
tradition. Basic scientists in pharmacology and physiology tend to use the
F test. Scientists in fields like ecology and population biology tend to use
AIC approach.

If the models are not nested, then the F test is not valid, so you should
choose the information theory approach. Note that Prism does not test
whether the models are nested.

How do these methods work to compare data sets?

The Compare tab of Prism lets you ask "Do the best-fit values of
selected unshared parameters differ between data sets?" or "Does one
curve adequately fit all data sets?". Applying the F test or Akaike's
method to answering these questions is straightforward. Prism compares
the sum-of-squares of two fits.

• In one fit, the model is separately fit to each data set, and the
goodness-of-fit is quantified with a sum-of-squares. The sum of these
sum-of-square values quantifies the goodness of fit of the family of
curves fit to all the data sets.

• The other fit is a global fit to all the data sets at once, sharing specified
parameters. If you ask Prism whether one curve adequately fits all data
sets, then it shares all the parameters.

These two fits are nested (the second is a simpler case of the first, with
fewer parameters to fit) so the sums-of-squares (actually the sum of
sum of squares for the first fits) can be compared using either the F
test or Akaike's method.

Don't compare curves using $R^2$ or the adjusted $R^2$

$R^2$ is a measure of how well a model fits your data, so it would seem to
make sense to choose among competing models by picking the one that
has the lowest $R^2$ or adjusted $R^2$. In fact, it works really poorly (1). Don't
use this method!

Interpreting comparison of models

2.6.3 How the F test works to compare models

The extra sum-of-squares F test compares the fits of two nested models. Nested means one model (the simpler one, model 1 below) is a special case of the other model (the more complicated one; model 2 below).

If the simpler model is correct, the relative increase in the sum of squares (going from more complicated to simpler model) is expected to equal the relative increase in degrees of freedom. In other words, if the simpler model is correct you expect that:

\[
\frac{SS_1 - SS_2}{SS_2} \approx \frac{DF_1 - DF_2}{DF_2}
\]

SS1 is the sum-of-squares for the simpler model (which will be higher) and SS2 is the sum-of-squares of the more complicated model. Recall that the number of degrees of freedom equals the number of data points minus the number of parameters. The more complicated model has more parameters so has fewer degrees of freedom.

If the more complicated model is correct, then you expect the relative increase in sum-of-squares (going from complicated to simple model) to be greater than the relative increase in degrees of freedom:

\[
\frac{SS_1 - SS_2}{SS_2} > \frac{DF_1 - DF_2}{DF_2}
\]

The F ratio quantifies the relationship between the relative increase in sum-of-squares and the relative increase in degrees of freedom.

\[
F = \frac{(SS_1 - SS_2)/SS_2}{(DF_1 - DF_2)/DF_2}
\]

That equation is more commonly shown in an equivalent form:

\[
F = \frac{(SS_1 - SS_2)/(DF_1 - DF_2)}{SS_2/DF_2}
\]
F ratios are always associated with a certain number of degrees of freedom for the numerator and a certain number of degrees of freedom for the denominator. This F ratio has DF1-DF2 degrees of freedom for the numerator, and DF2 degrees of freedom for the denominator.

If the simpler model is correct you expect to get an F ratio near 1.0. If the ratio is much greater than 1.0, there are two possibilities:

- The more complicated model is correct.
- The simpler model is correct, but random scatter led the more complicated model to fit better. The P value tells you how rare this coincidence would be.

The P value answers this question:

If model 1 is really correct, what is the chance that you would randomly obtain data that fits model 2 so much better?

If the P value is low, conclude that model 2 is statistically significantly better than model 1. Otherwise, conclude that there is no compelling evidence supporting model 2, so accept the simpler model (model 1).

### 2.6.4 How the AICc computations work

While the theoretical basis of Akaike's method is difficult to follow, it is easy to do the computations and make sense of the results.

The fit of any model to a data set can be summarized by an information criterion developed by Akaike, called the AIC. If you accept the usual assumptions of nonlinear regression (that the scatter of points around the curve follows a Gaussian distribution), the AIC is defined by a simple equation from the sum-of-squares and number of degrees of freedom of the two models. It is not possible to make sense of this AIC value itself, because its units depend on which units you use for your data.

To compare models, it is only the difference between the two AIC values that matter. When you take the difference, the units cancel out and the result is unitless.
\[ \Delta \text{AIC} = N \times \ln \left( \frac{SS_2}{SS_1} \right) + 2\Delta DF \]

The equation now makes intuitive sense. Like the F test, it balances the change in goodness-of-fit as assessed by sum-of-squares with the change in the number of degrees of freedom (due to differences in the number of parameters to be fit). Since model 1 is the simpler model, it will almost always fit worse, so SS1 will be greater than SS2. Since the logarithm of a fraction is always negative, the first term will be negative. Model 1 has fewer parameters and so has more degrees of freedom, making the last term positive. If the net result is negative, that means that the difference in sum-of-squares is more than expected based on the difference in number of parameters, so you conclude that the more complicated model is more likely.

Prism reports the difference between the two AICc values as the AICc of the simpler model minus the AICc of the more complicated model. When the more complicated (more parameters) model has the lower AICc and so is preferred, Prism reports the difference of AICc as a positive number. When the simpler model has the lower AICc and so is preferred, Prism reports the difference of AICc as a negative number.

The equation above helps you get a sense of how AIC works – balancing change in goodness-of-fit vs. the difference in number of parameters. But you don’t have to use that equation. Just look at the individual AIC values, and choose the model with the smallest AIC value. That model is most likely to be correct.

Prism actually doesn't report the AIC, but rather the AICc. That value includes a correction for low sample size. The equation is a bit more complicated, and is more accurate with small sample size. With larger sample sizes, the AIC and AICc are almost the same.

Note that these calculations are based on information theory, and do not use the traditional “hypothesis testing” statistical paradigm. Therefore there is no P value, no conclusion about “statistical significance”, and no “rejection” of a model.

From the difference in AICc values, Prism calculates and reports the probability that each model is correct, with the probabilities summing to 100%. If one model is much more likely to be correct than the other
(say, 1% vs. 99%), you will want to choose it. If the difference in
likelihood is not very big (say, 40% vs. 60%), you will know that either
model might be correct, so will want to collect more data. These
probabilities are computed with this equation, where Δ is the difference
between the AICc values.

\[
\text{probability} = \frac{e^{0.5\Delta}}{1 + e^{0.5\Delta}}
\]

Note that this method simply compares the fits of the two models you
chose. It is possible that a third model, one you didn't choose, fits far
better than either model you chose.

2.7 Outlier elimination and robust nonlinear regression

Nonlinear regression, like linear regression, assumes that the
scatter of data around the ideal curve follows a Gaussian or
normal distribution. Outliers can violate this assumption and
invalidate nonlinear regression results. To deal with outliers,
Prism offers robust regression and automatic outlier removal.

- **What are the advantages of removing outliers.**
  But isn't it cheating?

- Outlier removal has to be used carefully. It must be avoided in many situations.

- Once you've identified an outlier, beware or removing it from graphs and analyses. An outlier is not always a 'bad' point.

- Prism uses an outlier removal method created by GraphPad Software --the ROUT method. Learn how it works.

- When identifying outliers, the step is to fit a curve using a robust method designed so outliers won't affect the curve much. Learn about robust nonlinear regression.
2.7.1 When to use automatic outlier removal

The problem with outliers

Nonlinear regression, like linear regression, assumes that the scatter of data around the ideal curve follows a Gaussian or normal distribution. This assumption leads to the familiar goal of regression: to minimize the sum of the squares of the vertical or Y-value distances between the points and the curve. However, experimental mistakes can lead to erroneous values – outliers. Even a single outlier can dominate the sum-of-the-squares calculation, and lead to misleading results.

Is it 'cheating' to remove outliers?

Some people feel that removing outliers is ‘cheating’. It can be viewed that way when outliers are removed in an \textit{ad hoc} manner, especially when you remove only outliers that get in the way of obtaining results you like. But leaving outliers in the data you analyze is also ‘cheating’, as it can lead to invalid results.

Here is a Bayesian way to think about systematic approaches to removing outliers. When a value is flagged as an outlier, there are two possibilities.

- A coincidence occurred, the kind of coincidence that happens in few percent of experiments even if the entire scatter is Gaussian (depending on how aggressively you define an outlier).
- A ‘bad’ point got included in your data.

Which possibility is more likely?

It depends on your experimental system.

If your experimental system generates a ‘bad’ point in a few percent of experiments, then it makes sense to eliminate the point as an outlier. It is more likely to be a ‘bad’ point than a ‘good’ point that just happened to be far from the curve.

If your system is very pure and controlled, so ‘bad’ points occur very rarely, then it is more likely that the point is far from the curve due to chance (and not mistake) and you should leave it in. Alternatively in that
case, you could set Q to a lower value in order to only detect outliers that are much further away.

2.7.2 When to avoid automatic outlier removal

**Outlier elimination is misleading when you are fitting the wrong model**

The left panel above shows the data fit to a dose response curve. In this figure, one of the points is a significant outlier. But this interpretation assumes that you’ve chosen the correct model. The right panel shows the data fit to an alternative bell-shaped dose-response model, where high doses evoke a smaller response than does a moderate dose. The data fit this model very well, with no outliers detected (or even suspected).

This example points out that outlier elimination is only appropriate when you are sure that you are fitting the correct model.

**Outlier elimination is misleading when data points are not independent**

The left panel above show data fit to a dose-response model with one point (in the upper right) detected as an outlier. The right panel shows...
that the data really come from two different experiments. Both the lower and upper plateaus of the second experiment (shown with upward pointing triangles) are higher than those in the first experiment (downward pointing triangles). Because these are two different experiments, the assumption of independence was violated in the analysis in the left panel. When we fit each experimental run separately, no outliers are detected.

**Outlier elimination is misleading when you chose incorrect weighting factors**

![Graph showing outlier elimination](image)

The left panel above shows data fit to a dose-response model. Four outliers were identified (two are almost superimposed). But note that the values with larger responses (Y values) also, on average, are further from the curve. This makes least-squares regression inappropriate. To account for the fact that the SD of the residuals is proportional to the height of the curve, we need to use weighted regression. The right panel shows the same data fit to the same dose-response model, but minimizing sum of the squares of the distance of the point from the curve divided by the height of the curve, using relative weighting. Now no outliers are identified. Using the wrong weighting method created false outliers.

### 2.7.3 Outliers aren't always 'bad' points

**Definition of an 'outlier'**
The term 'outlier' is defined fairly vaguely, but refers to a value that is far from the others. In Prism's nonlinear regression, an outlier is a point that is far from the best-fit curve defined by robust regression.

Of course, there is some possibility that an outlier really comes from the same Gaussian population as the others, and just happens to be very high or low. You can set the value of $Q$ to control how aggressively Prism defines outliers.

**Outliers are not always due to mistakes**

Nonlinear regression is usually used with experimental data, where X is a variable like time or concentration or some other variable you manipulate in the experiment. Since all the scatter is due to experimental error, it can make sense to eliminate any extreme outlier since it is almost certainly the result of an experimental mistake.

In other situations, each data point can represent a different individual. In this case, an outlier may not be due to experimental mistakes, but rather be the result of biological variation, or differences in some other variable that is not included in your model. Here, the presence of the outlier may be the most interesting finding in the study. While the ROUT outlier method might prove useful to flag an outlier in this situation, it would be a big mistake to automatically exclude such outliers without further thought (or experimentation).

In quality control analyses, an outlier can tell you about a process that is out of control. You wouldn't want to delete outliers, without first figuring out why the value is far from the others. The outlier might be telling you something important.

### 2.7.4 The ROUT method of identifying outliers

**How the ROUT method of removing outliers works**

Prism offers a unique approach to identifying and removing outliers, detailed in reference 1. Because this method combines robust regression and outlier removal, we call it the **ROUT** method.

The ROUT method of regression follows these steps.
1. Our robust nonlinear regression method is used to fit a curve that is not influenced by outliers.

2. The residuals of the robust fit are analyzed to identify any outliers. This step uses a new outlier test adapted from the False Discovery Rate approach of testing for multiple comparisons.

3. Remove the outliers, and perform ordinary least-squares regression on the remaining data.

How to use the ROUT method with Prism

Although the ROUT method requires three steps (listed above), Prism does all this automatically. All you have to do is check an option on the Fit tab of nonlinear regression dialog:

Prism then identifies the outliers, eliminates them, and fits the remaining points. The outliers are shown in a separate table, and the number of outliers is tabulated on the main results table.

The ROUT coefficient Q

The value of Q determines how aggressively the ROUT method defines outliers. The mathematical details are explained in reference 1. This value is set in the Weights tab of the Nonlinear regression dialog.

If you set Q to a higher value, the threshold for defining outliers is less strict. This means that Prism will have more power to detect outliers, but also will falsely detect 'outliers' more often.

If you set Q to a lower value, the threshold for defining outliers is stricter. This means that Prism will have a less power to detect real outliers, but also have a smaller chance of falsely defining a point to be an outlier.

Unless you have a strong reason to choose otherwise, we recommend sticking with the default value of 1%. Our simulations have shown that if all the scatter is Gaussian, Prism will falsely find one or more outliers in about 2-3% of experiments. This does not mean that a few percent of all
values are declared to be outliers, but rather that one or more outliers will be detected in a few percent of experiments. If there really are outliers present in the data, Prism will detect them with a False Discovery Rate less than 1%.

**Unequal weighting, robust regression and outlier removal**

As we explain in reference 1, unequal weighting is not useful with robust regression. The problem is that outliers can get way too much weight.

If you choose both unequal weighting and robust fitting, therefore, Prism does the fitting assuming equal weights. However it does use your weighting choice when creating a table of residuals.

If you choose both unequal weighting and automatic outlier removal, Prism first fits using robust regression (ignoring your weighting choice). It does use the weighting factors when identifying the outliers, as explained in reference 1. It then performs weighted nonlinear regression on the outlier-depleted data.


### 2.7.5 Robust nonlinear regression

**The need for robust regression**

Nonlinear regression, like linear regression, assumes that the scatter of data around the ideal curve follows a Gaussian or normal distribution. This assumption leads to the familiar goal of regression: to minimize the sum of the squares of the vertical or Y-value distances between the points and the curve. This standard method for performing nonlinear (or linear regression) is called **least-squares**.
Experimental mistakes can lead to erroneous values whose values are way too high or too low—outliers. Even a single outlier can dominate the sum-of-the-squares calculation, and lead to misleading results. One way to cope with this problem is to perform a robust fit using a method that is not very sensitive to violations of the Gaussian assumption. Another approach is to use automatic outlier elimination to identify and remove the outliers, and then run least-squares regression. Prism offers both choices. The outlier identification method actually first does a robust fit, so it has a baseline from which to decide when a point is too far from that baseline so becomes defined as an outlier. Then after removing those outliers, it performs a standard least-squares fit on the remaining points.

**How robust regression works**

Based on a suggestion in Numerical Recipes (1), we based our robust fitting method on the assumption that variation around the curve follows a Lorentzian distribution, rather than a Gaussian distribution. Both distributions are part of a family of t distributions:

- df=1 (Lorentzian or Cauchy)
- df=4
- df=10
- df=∞ (Gaussian)

The widest distribution in that figure, the t distribution for df=1, is also known as the Lorentzian distribution or Cauchy distribution. The Lorentzian distribution has wide tails, so outliers are fairly common and therefore have little impact on the fit.

We adapted the Marquardt nonlinear regression algorithm to accommodate the assumption of a Lorentzian (rather than Gaussian) distribution of residuals, and explain the details in reference 2.
When does it make sense to choose robust nonlinear regression?

The main use of robust regression in Prism is as a 'baseline' from which to remove outliers. You may want to experiment with robust regression in order to better understand the outlier-removal method (which begins with robust regression). You may also find it useful if your only goal is to interpolate from a standard curve, and that standard curve has one or more outliers.

We recommend that you avoid robust regression (at least as implemented by Prism) for most uses, because it has these drawbacks:

- Robust regression cannot calculate standard errors or confidence intervals for the parameters.
- Robust regression cannot plot confidence or prediction bands.
- Robust regression cannot compute the degree to which parameters are intertwined. It cannot compute the covariance matrix or the dependency values.
- Robust regression cannot compare the fits of two models or two data sets.

The inability to compare models does not matter in most contexts. But the inability to provide diagnostic information (SE, CI, Covariance matrix, Dependency, Confidence and prediction bands) seriously limits the usefulness of robust regression for most research.

References

2.8 How nonlinear regression works

If you want to really understand how nonlinear regression works, you will need to learn matrix algebra and study advanced math books. But you can use nonlinear regression effectively so long as you understand the basic principles of regression.

- **Why minimize the sum-of-squares?**

- **How nonlinear regression works**

- **The need for unequal weighting in nonlinear regression**

- **How standard errors and confidence intervals are computed**

- **How confidence and prediction bands are computed**

- **Replicates**

- **How dependency is calculated**

2.8.1 Why minimize the sum-of-squares?

The goal of nonlinear regression is to adjust the values of the model's parameters to find the curve that best predicts $Y$ from $X$. More precisely, the goal of regression is to minimize the sum of the squares of the vertical distances of the points from the curve.

Why minimize the sum of the squares of the distances? Why not simply minimize the sum of the actual distances?

If the random scatter follows a Gaussian distribution, it is far more likely to have two medium size deviations (say 5 units each) than to have one small deviation (1 unit) and one large (9 units). A procedure that minimized the sum of the absolute value of the distances would have no preference over a curve that was 5 units away from two points and one that was 1 unit away from one point and 9 units from another. The sum of the distances (more precisely, the sum of the absolute
value of the distances) is 10 units in each case. A procedure that minimizes the sum of the squares of the distances prefers to be 5 units away from two points (sum-of-squares = 50) rather than 1 unit away from one point and 9 units away from another (sum-of-squares = 82). If the scatter is Gaussian (or nearly so), the curve determined by minimizing the sum-of-squares is most likely to be correct.

### 2.8.2 How nonlinear regression works

#### The basic idea of nonlinear regression

You won’t be able to understand the mathematical details of nonlinear regression unless you first master matrix algebra. But the basic idea is pretty easy to understand. Every nonlinear regression method follows these steps:

1. Start with initial estimated values for each parameter in the equation.

2. Generate the curve defined by the initial values. Calculate the sum-of-squares -- the sum of the squares of the vertical distances of the points from the curve. (Or compute the weighted sum-of-squares if you are including weighting factors.)

3. Adjust the parameters to make the curve come closer to the data points -- to reduce the sum-of-squares. There are several algorithms for adjusting the parameters, as explained below.

4. Adjust the parameters again so that the curve comes even closer to the points. Repeat.

5. Stop the calculations when the adjustments make virtually no difference in the sum-of-squares.

6. Report the best-fit results. The precise values you obtain will depend in part on the initial values chosen in step 1 and the stopping criteria of step 5. This means that repeat analyses of the same data will not always give exactly the same results.

#### The Marquardt method

Step 3 is the only difficult one. Prism (and most other nonlinear regression programs) uses the method of Marquardt and Levenberg,
which blends two other methods, the method of linear descent and the method of Gauss-Newton. The best way to understand these methods is to follow an example. Here are some data to be fit to a typical binding curve (rectangular hyperbola).

![Graph showing specific binding vs ligand concentration](image)

You want to fit a binding curve to determine Bmax and Kd using the equation

\[ Y = \frac{B_{\text{max}} \cdot X}{K_d + X} \]

How can you find the values of Bmax and Kd that fit the data best? You can generate an infinite number of curves by varying Bmax and Kd. For each of the generated curves, you can compute the sum-of-squares to assess how well that curve fits the data. The following graph illustrates the situation.
The X- and Y-axes correspond to two parameters to be fit by nonlinear regression (Bmax and Kd in this example). The Z-axis is the sum-of-squares. Each point on the surface corresponds to one possible curve. The goal of nonlinear regression is to find the values of Bmax and Kd that make the sum-of-squares as small as possible (to find the bottom of the valley).

The method of linear descent follows a very simple strategy. Starting from the initial values, try increasing each parameter a small amount. If the sum-of-squares goes down, continue. If the sum-of-squares goes up, go back and decrease the value of the parameter instead. You've taken a step down the surface. Repeat many times. Each step will usually reduce the sum-of-squares. If the sum-of-squares goes up instead, the step must have been so large that you went past the bottom and back up the other side. If this happens, go back and take a smaller step. After repeating these steps many times, you will reach the bottom.

The Gauss-Newton method is a bit harder to understand. As with the method of linear descent, start by computing how much the sum-of-squares changes when you make a small change in the value of each parameter. This tells you the slope of the sum-of-squares surface at the point defined by the initial values. If the equation really is linear, this is enough information to determine the shape of the entire sum-of-squares surface, and thus calculate the best-fit values of Bmax and Kd in one step. With a linear equation, knowing the slope at one point tells you
everything you need to know about the surface, and you can find the minimum in one step. With nonlinear equations, the Gauss-Newton method won't find the best-fit values in one step, but that step usually improves the fit. After repeating many iterations, you reach the bottom.

This method of linear descent tends to work well for early iterations, but works slowly when it gets close to the best-fit values (and the surface is nearly flat). In contrast, the Gauss-Newton method tends to work badly in early iterations, but works very well in later iterations. The two methods are blended in the method of Marquardt (also called the Levenberg-Marquardt method). It uses the method of linear descent in early iterations and then gradually switches to the Gauss-Newton approach.

Prism, like most programs, uses the Marquardt method for performing nonlinear regression.

### References


#### 2.8.3 Nonlinear regression with unequal weights

**How Prism implements weighting**

Prism 5 and 6 weight by the Y value of the curve. Prism 4 and earlier weighted by the Y value of the data. The distinction is subtle and rarely matters much, but our simulations show that the results are sometimes more accurate when weights are based on the value of the curve rather than the data.

The situation is a bit tricky. The goal is to adjust the values of the parameters to minimize the weighted sum-of-squares. But the values of the weights depend on the values of those parameters. Here is how Prism 6 resolves this issue:

1. Perform the first iteration without any weights. If the initial parameter values are not very accurate, weighted regression can cause problems on the first iteration. This step was introduced in
Prism 5.02 and 5.0b. Without it, some fits with bad initial values simply won't converge.

2. Compute Ycurve at all values of X based on the curve generated by this first iteration of nonlinear regression, and use those values to compute weights for each points.

3. Keeping those weights constant, use nonlinear regression to minimize the weighted sum-of-squares.

4. Recompute Ycurve based on the results of that curve fit and generate a new set of weights.

5. Keeping those new weights constant, use nonlinear regression again to minimize the weighted sum-of-squares.

6. Repeat steps 4 and 5 until the weighted sum-of-squares no longer changes.

Weighting with robust regression or automatic outlier removal

As we explain in reference 1, it doesn't make sense to perform robust regression using unequal weights. The problem is that outliers can get too much weight.

If you choose both unequal weighting and robust fitting, therefore, Prism does the fitting assuming equal weights. However it uses your weighting choice when creating a table of residuals, and when counting the number of outliers (a choice you can make in the preferences tab).

If you choose both unequal weighting and automatic outlier removal, Prism first fits using robust regression (ignoring your weighting choice), and then uses the weighting factors in identifying the outliers, as explained in reference 1.

Reference


2.8.4 How standard errors are computed

Prism uses a standard method to compute the standard error and
confidence interval for each parameter fit with nonlinear regression.

Each parameter's standard error is computed using this equation:

\[ \text{SE}(P_i) = \sqrt{\left(\frac{SS}{DF}\right) \cdot \text{Cov}(i,i)} \]

where:

- \( P_i \) : i-th adjustable (non-constant) parameter
- \( SS \) : sum of squared residuals
- \( DF \) : degrees of freedom (the number of data points minus number of parameters fit by regression)
- \( \text{Cov}(i,i) \) : i-th diagonal element of covariance matrix
- \( \sqrt() \) : square root

### 2.8.5 How confidence intervals are computed

**Asymptotic confidence intervals of parameters**

Asymptotic confidence intervals are always centered on the best-fit value of the parameter, and extend the same distance above and below that value.

The 95% confidence intervals are computed by this equation:

\[ \text{From } [\text{BestFit} - t \times \text{SE}] \text{ TO } [\text{BestFit} + t \times \text{SE}] \]

where BestFit is the best fit value for the parameter, SE is its standard error, and \( t \) is the value from the t distribution for the desired level of confidence (95% is standard) and the number of degrees of freedom (which equals the number of data points minus the number of parameters fit by regression). With 95% confidence and many degrees of freedom (more than a few dozen), this multiplier is very close to 1.96. Note that the value of \( t \) is not computed from your data, but is a constant that depends on the confidence level you choose, the number of data points, and the number of parameters.
Profile likelihood asymmetrical confidence intervals (new to Prism 7)

Until Prism 7, Prism only reported asymptotic confidence intervals that are always symmetrical around the best-fit values.

For some parameters in some models, an asymmetrical interval does a much better job of expressing precision. Prism 7 now offers this with profile likelihood confidence intervals, as a choice in the Confidence tab of the nonlinear regression dialog. The disadvantages are that they are unfamiliar to many and that they take longer to compute (but with a fast computer, you may not even notice unless you have huge data sets and choose a user-defined equation).

The idea is pretty simple. The extra sum-of-squares test compares two models.

- The more complicated model is the model you chose. The entire page of results is for this model.

- The simpler model fixes one parameter to a constant value. The idea is to fix the parameter to various values until you find the right value (as explained below).

Here is a very simplified algorithm that sort of explains the idea behind the method. Define the sum-of-squares to be SS and the degrees of freedom to be DF.

1. Set a variable Delta to the SE of the parameter.

2. Hold the parameter fixed to its best-fit value minus delta, and run the fit again letting all the other parameter values vary. Record the new SS and DF of this fit.

3. Compare the original best-fit with this fit that forces the parameter to be decreased by delta using the extra sum-of-squares F test. The second fit holds one parameter to a constant value so fits one fewer parameter as so has one more degree of freedom. Compute the P value.

   a. If the P value is less than 0.05, delta is too large. Make it smaller and go back to step 2.
b. If the P value is greater than 0.05, delta is too small. Make it larger and go back to step 2.

c. If the P value is very close to 0.05, then the lower confidence limit equals the original best fit value minus delta.

4. Holding your parameter fixed to its best-fit value plus delta, run the fit again letting all the other parameter values vary. Record the SS and DF of this fit.

5. Compare the original best-fit with this fit that forces the parameter to be increased by delta using the extra sum-of-squares F test. Compute the P value.

a. If the P value is less than 0.05, delta is too large. Make it smaller and go back to step 5.

b. If the P value is greater than 0.05, delta is too small. Make it larger and go back to step 5.

c. If the P value is very close to 0.05, then the upper confidence limit equals the original best fit value plus delta.

6. Repeat for each parameter.

Prism actually uses the steps detailed by Venzon and Moolgavkar(1) for each parameter. This method creates a likelihood profile for each parameter. For various possible values of the parameter, the algorithm fits the curve (optimizing the other parameters) and determines the likelihood that the data would have come from this model. The confidence interval is the range of parameter values where the likelihood is not too much lower than its maximum value. Of course, "too low" is defined rigorously.

The maximum likelihood is at the best-fit value of the parameter. When these profiles are graphed in texts, it is usually the negative logarithm of the likelihood that is plotted. The maximum likelihood is the same as the minimum -log(likelihood) so in these graphs the best fit value is the X value where Y is at its lowest value.

If you assume all residuals follow a Gaussian distribution, maximizing the likelihood is the same as minimizing the sum-of-squares.
Notes:

- The final delta value for computing the upper confidence limit may not be equal to (or even be close to) the final value of delta for computing the lower limit. That's why the confidence interval may be asymmetrical around the best-fit value.

- The P value goal of 0.05 above is used only when you want 95% confidence intervals. If you want 99% confidence intervals, use 0.01, etc.

- The method in reference 1 is way more clever than described above so takes fewer computations.

- The confidence intervals computed this way are for just that one parameter. We expect that each confidence interval has a 95% chance of including the true parameter value. The 95% does not apply to the set of intervals. It is not correct to say that we expect there to be a 95% chance that all the confidence intervals include the respective true parameter values.

- When computing the extra sum of squares F test above, note that the two models differ by one degree of freedom. This is because we are fixing one parameter and letting Prism fit the others. Some publications (2) assume that you are fixing all the parameters, not just one. So the two models entered into the F test differ by K degrees of freedom, where K is the number of parameters fit. These intervals are wider, and I think the intent is that the 95% confidence level apply simultaneously to all the intervals, rather than to just one. Prism does not use this approach. With prism, the two models being compared always differ by one degree of freedom.

- The method we use is also described by Watts(3). Prism matches the results he presents in Table IV for data in Table III.

- In some cases, the method is unable to find one of the confidence limits and reports "???" instead of a value.

**The confidence interval for the difference or ratio of two parameters**

When defining an equation, you can choose (in the Transforms to Report tab) to have Prism also report the difference or ratio (etc) of two
parameters, with standard error and confidence interval. The calculations account for the covariance of the two parameters. See this document.

Reference


2.8.6 How confidence and prediction bands are computed

The difference between confidence and prediction bands

Note the difference between confidence and prediction bands:

- The 95% confidence bands enclose the area that you can be 95% sure contains the true curve. If you have many data points, the confidence bands will be near the line or curve, and most of your data will lie outside the confidence bands.

- The 95% prediction bands enclose the area that you expect to enclose 95% of future data points. They are wider than confidence bands -- much wider with large data sets.

Also distinguish between the 95% confidence interval of the parameters (a range of values), and the 95% confidence bands around the curve.

How confidence and prediction bands are computed

The calculation of the confidence and prediction bands are fairly standard, and can only be expressed with matrices. A brief explanation follows. More details can be found here.

First, define $G|x$, which is the gradient of the parameters at a particular value of $X$ and using all the best-fit values of the parameters. The result
is a vector, with one element per parameter. For each parameter, it is defined as $\frac{dY}{dP}$, where Y is the Y value of the curve given the particular value of X and all the best-fit parameter values, and P is one of the parameters.)

$G'|x$ is that gradient vector transposed, so it is a column rather than a row of values.

Cov is the covariance matrix (inverted Hessian from last iteration). It is a square matrix with the number of rows and columns equal to the number of parameters. Each item in the matrix is the covariance between two parameters. Note that this is the actual covariance matrix, which is distinct from the normalized covariance matrix (where each value is between -1 and 1) that Prism can report.

Now compute $c = G|x * Cov * G'|x$. The result is a single number for any value of X.

The confidence and prediction bands are centered on the best fit curve, and extend above and below the curve an equal amount.

The confidence bands extend above and below the curve by:

$$= \sqrt{c} \sqrt{SS/DF} \times \text{CriticalT(Confidence\%}, \ DF)$$

The prediction bands extend a further distance above and below the curve, equal to:

$$= \sqrt{c+1} \sqrt{SS/DF} \times \text{CriticalT(Confidence\%}, \ DF)$$

In both these equations, the value of $c$ (defined above) depends on the value of X, so the confidence and prediction bands are not a constant distance from the curve. The value of SS is the sum-of-squares for the fit, and DF is the number of degrees of freedom (number of data points minus number of parameters). CriticalT is a constant from the t distribution based on the amount of confidence you want and the number of degrees of freedom. For 95% limits, and a fairly large df, this value is close to 1.96. If DF is small, this value is higher.

2.8.7 Replicates

If you entered data with replicates (in side-by-side subcolumns), Prism gives you two choice for how to deal with the replicates.
• Consider each replicate as an individual point.

• Fit the means of each set of replicates.

The rest of this page explains how to decide. When in doubt, choose to fit individual replicates. The other choice is only rarely useful.

**Independent replicates**

In most experiments, it is fair to consider each replicate to be an independent data point. Each particular replicate is subject to random factors, which may increase or decrease its value. Each random factor affects individual replicates, and no random factor affects the replicates as a group. In any kind of biochemical experiment, where each value comes from a test tube or plate well, the replicates are almost certain to be independent.

When your replicates are independent, Prism will treat each replicate as a separate point. If there are four replicates at one X value and two at another, the four replicates will automatically get twice the weight, since the program considers them to be four separate data points.

If you ask Prism to fit the mean values, rather than individual replicates, you won't get valid standard errors and confidence intervals. If you have different number of replicates at different X values, you will lose the extra weights that the points with more replicates deserve, so will get incorrect best-fit values.

**Replicates that are not independent**

In some experimental situations, the replicates are not independent. Random factors can affect all the replicates at once. Two examples:

• You performed a binding experiment with a single tube at each concentration, but measured the radioactivity in each tube three times. Those three values are not independent. Any experimental error while conducting the experiment would affect all the replicates.

• You performed a dose-response experiment, using a different animal at each dose with triplicate measurements. The three measurements are not independent. If one animal happens to respond more than the others, that will affect all the replicates. The replicates are not independent.
Treating each replicate as a separate data point would not be appropriate in these situations. Most of the random variation is between tubes (first example) or animals (second example). Collecting multiple replicates does not give you much additional information. Certainly, each replicate does not give independent information about the values of the parameters. Here is one way to look at this. Imagine that you have performed a dose-response experiment with a separate animal for each dose. You measure one animal in duplicate (for one dose) and another animal (another dose) ten times. It would be a mistake to enter those as individual values, because that would give five times more weight to the second dose compared to the first. The random factors tend to affect the animal, not the measurement, so measuring an animal ten times does not give you five times more information about the true value than measuring it two times.

Since each tube (first example, above) or animal (second example) is the experimental unit, you should enter each tube or animal once. If you measured several replicates, average these and enter the average. Don’t enter individual values. Don’t weight the means by sample size. Doing so would inflate the number of degrees of freedom inappropriately, and give you SE that are too small and CI that are too narrow. Doing so, when you have unequal number of replicates would give artificial, and undeserved, weight to the tubes or animals with more replicates, so would affect the best-fit curve and you would get less than optimal best fit parameter values.

If you are confused by the choice of fitting to individual replicates vs. the mean, choose to fit individual replicates (which is the default

### 2.8.8 How dependency is calculated

#### What is dependency?

When the model has two or more parameters, as is almost always the case, the parameters can be intertwined.

What does it mean for parameters to be intertwined? After fitting a model, change the value of one parameter but leave the others alone. The curve moves away from the points. Now, try to bring the curve back so it is close to the points by changing the other parameter(s). If you can bring the curve closer to the points, the parameters are intertwined. If you can bring the curve back to its original position, then the parameters are redundant.
Prism can quantify the relationships between parameters by reporting the correlation matrix or reporting dependency.

**Interpreting dependency**

You can interpret dependency without knowing much about how it is calculated. Read on if you are interested in knowing how the value is computed.

**Example of dependency calculations**

This example is an exponential decay (taken from pages 128-130 of the MLAB Applications Manual, www.civilized.com).

<table>
<thead>
<tr>
<th>Time</th>
<th>Signal</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.0</td>
<td>39.814</td>
</tr>
<tr>
<td>2.0</td>
<td>32.269</td>
</tr>
<tr>
<td>3.0</td>
<td>29.431</td>
</tr>
<tr>
<td>4.0</td>
<td>27.481</td>
</tr>
<tr>
<td>5.0</td>
<td>26.086</td>
</tr>
<tr>
<td>6.0</td>
<td>25.757</td>
</tr>
<tr>
<td>7.0</td>
<td>24.932</td>
</tr>
<tr>
<td>8.0</td>
<td>23.928</td>
</tr>
<tr>
<td>9.0</td>
<td>22.415</td>
</tr>
<tr>
<td>10.0</td>
<td>22.548</td>
</tr>
<tr>
<td>11.0</td>
<td>21.900</td>
</tr>
<tr>
<td>12.0</td>
<td>20.527</td>
</tr>
<tr>
<td>13.0</td>
<td>20.695</td>
</tr>
<tr>
<td>14.0</td>
<td>20.105</td>
</tr>
<tr>
<td>15.0</td>
<td>19.516</td>
</tr>
<tr>
<td>16.0</td>
<td>19.640</td>
</tr>
<tr>
<td>17.0</td>
<td>19.346</td>
</tr>
<tr>
<td>18.0</td>
<td>18.927</td>
</tr>
<tr>
<td>19.0</td>
<td>18.857</td>
</tr>
<tr>
<td>20.0</td>
<td>17.652</td>
</tr>
</tbody>
</table>

We will focus on the rate constant, K. The best fit value is 0.2149 sec⁻¹, which corresponds to a half-life of 3.225 seconds. Its SE is 0.0248 sec⁻¹, which corresponds to a 95% confidence interval of 0.1625 to 0.2674 sec⁻¹.

It is clear that the three parameters are not entirely independent. If you forced $K$ to have a higher value (faster decay), the curve would get further from the points. But you could compensate a bit by starting the
curve at a higher value and ending at a lower one (increase Span and decrease Plateau). The SE values of the parameters depend on one another.

Fix Span and Plateau to their best fit values, and ask Prism to fit only the rate constant K. This will not change the best fit value, of course, since we fixed Span and Plateau to their best-fit values. But the SE of K is lower now, equal to 0.008605. This makes sense. Changing the value of K has a bigger impact on goodness-of-fit (sum-of-squares) when you fix the Span and Plateau than it does when you allow the values of Span and Plateau to also change to compensate for the change in K.

The lower value of the SE of K when you fix the other parameters tells you that the uncertainty in K is dependent on the other parameters. We want to quantify this by computing the dependency.

Before we can compare the two SE values, we have to correct for a minor problem. When computing the SE, the program divides by the square root of the number of degrees of freedom (df). For each fit, df equals the number of data points minus the number of parameters fit by the regression. For the full fit, df therefore equals 20 (number of data points) minus 3 (number of parameters) or 17. When we held the values of Plateau and Span constant, there was only one parameter, so df=19. Because the df are not equal, the two SE values are not quite comparable. The SE when other parameters were fixed is artificially low. This is easy to fix. Multiply the SE reported when two of the parameters were constrained by the square root of 19/17. This corrected SE equals 0.00910.

Now we can compute the dependency. It equals 1.0 minus the square of the ratio of the two (corrected) SE values. So the dependency for this example equals 1.0-(0.0091/0.0248)², or 0.866. Essentially, this means that 86.6% of the variance in K is due to its interaction with other parameters.

Each parameter has a distinct dependency (unless there are only two parameters). The dependency of Span is 0.613 and the dependency of Plateau is 0.813.

**The origin of the idea of dependency**

There appears to be no paper to cite regarding the first use of dependency. The idea of dependency apparently was developed by Dick
Shrager at the NIH, and then enhanced by Gary Knott. MLAB was the first software to compute dependency, and it is explained well in the MLAB manual. GraphPad Prism simply implements the method as it is explained there. (I learned this history in an email from Gary Knott in 2007). Here is an early paper that discusses the basic ideas of dependency, but it is defined differently (ranging from 1 to infinity, rather than 0 to 1, and the math is not fully explained.

2.8.9 Who developed nonlinear regression?

According to wikipedia (Sept. 2011), the algorithm was first published by Kenneth Levenberg, while working at the Frankford Army Arsenal. It was rediscovered by Donald Marquardt who worked as a statistician at DuPont and independently by Girard, Wynn and Morrison.


3 REGRESSION WITH PRISM 7

Prism offers several analyses that can fit lines and curves.

- Linear regression
- Deming linear regression
- Nonlinear regression (which also does polynomial regression)
- Spline and Lowess
- Interpolating from a standard curve (linear or nonlinear)
3.1 Linear regression with Prism

Linear regression fits a straight line through your data to find the best-fit value of the slope and intercept.

**How to: Linear regression**

- Finding the best-fit slope and intercept
- Interpolating from a linear standard curve
- Advice: When to fit a line with nonlinear regression

**Results of linear regression**

- Slope and intercept
- r², a measure of goodness-of-fit of linear regression
- Is the slope significantly different than zero?
- Comparing slopes and intercepts
- Runs test following linear regression
- Analysis checklist: Linear regression
- Graphing tips: Linear regression
- Questions and answers

**Deming regression**

- Key concepts: Deming regression
- How to: Deming regression
- Q&A: Deming Regression
- Analysis checklist: Deming regression

3.1.1 How to: Linear regression

3.1.1.1 Finding the best-fit slope and intercept

1. Create a data table

From the Welcome or New Table dialog, choose to create XY data table.

If you are just getting started, choose the sample data: Linear regression -- Compare slopes.
If you are entering your own data, choose the subcolumn format. Choose replicate values if you have replicates to enter. Prism can plot error bars automatically. You can also choose to enter data where the mean and SD (or SEM) have already been calculated. In this case, if you want to take into account variations in the SD from point to point, use nonlinear regression to fit the line.

2. Enter data

If you chose sample data, you'll see these values:

If you enter Y values for several data sets (column A, B and C), Prism will report regression results for X vs. YA, for X vs. YB, and for X vs. YC. It can also test whether the slopes (and intercepts) differ significantly.

If the different data sets don't share the same X values, use different rows for different data sets like this:
3. Analysis choices

Click Analyze, and then choose linear regression from the list of XY analyses.

**Force the line to go through a specified point (such as the origin)?**

If you choose regression, you may force the line to go through a particular point such as the origin. In this case, Prism will determine only the best-fit slope, as the intercept will be fixed. Use this option when scientific theory tells you that the line must go through a particular point (usually the origin, X=0, Y=0) and you only want to know the slope. This situation arises rarely.

Use common sense when making your decision. For example, consider a protein assay. You measure optical density (Y) for several known concentrations of protein in order to create a standard curve. You then want to interpolate unknown protein concentrations from that standard curve. When performing the assay, you adjusted the spectrophotometer so that it reads zero with zero protein. Therefore you might be tempted to force the regression line through the origin. But this constraint may result in a line that doesn't fit the data very well. Since you really care that the line fits the standards very well near the unknowns, you will probably get a better fit by not constraining the line.

If in doubt, you should let Prism find the best-fit line without any constraints.

**Fit linear regression to individual replicates or to means?**

If you collected replicate Y values at every value of X, there are two ways to calculate linear regression. You can treat each replicate as a separate point, or you can average the replicate Y values, to determine the mean Y value at each X, and do the linear regression calculations using the means.

You should consider each replicate a separate point when the sources of experimental error are the same for each data point. If one value happens to be a bit high, there is no reason to expect the other replicates to be high as well. The errors are independent.

Average the replicates and treat the mean as a single value when the replicates are not independent. For example, the replicates would not be
independent if they represent triplicate measurements from the same animal, with a different animal used at each value of X (dose). If one animal happens to respond more than the others, that will affect all the replicates. The replicates are not independent.

**Test departure from linearity with runs test**

See [Runs test](#).

**Test whether slope and intercept are significantly different**

If you have entered data for two or more datasets, Prism can test whether the slopes differ significantly.

**Confidence and prediction bands**

Learn about [confidence and prediction bands](#).

### 3.1.1.2 Interpolating from a linear standard curve

1. **Create a data table**

   From the Welcome or New Table dialog, choose to create an XY data table.

   If you are just getting started, choose the sample data: Linear regression -- Interpolate from standard curve

   If you are entering your own data, choose the subcolumn format. Choose replicate values if you have replicates to enter. Prism can plot error bars automatically. You can also choose to enter data where the mean and SD (or SEM) have already been calculated. In this case, if you want to take into account variations in the SD from point to point, use [nonlinear regression to fit the line](#).

2. **Enter data**

   Enter the unknowns below the standards on the same table. Enter Y values with no X values in those rows (example below), or X values with no Y values in those rows. Optionally enter row titles to label those unknowns.
3. View the graph

4. Analysis choices

Click the Analyze button and from the list of XY analyses choose: Interpolate a Standard Curve.

Alternatively, you can click the “Interpolate a standard curve” button right on top of the Analyze button.
Choose Line in the list of Standard curves to interpolate. Unless you have a good reason, you can leave the other choices on that dialog set to their default values.

5. Inspect the results

View the graph with the superimposed line.

The results will appear in several pages, including one with the interpolated values, which will be in the same units as your original data.
The second page of results tabulates the best-fit values of the parameters and much more. For this example, we aren't too interested in these results.

### 3.1.1.3 Advice: When to fit a line with nonlinear regression

#### Linear regression is a special case of nonlinear regression

Linear regression is just a simpler, special, case of nonlinear regression. The calculations are a bit easier (but that only matters to programmers). You can use Prism's nonlinear regression analysis to fit a straight-line model, and the results will be identical to linear regression.

#### How to convert a linear regression analysis to nonlinear regression

To convert linear regression to nonlinear regression, bring up the Parameters dialog for linear regression, and click "More Choices" at the bottom.

#### Nonlinear regression offers more options

Conceptually, linear regression is just a special case of nonlinear regression. But Prism offers many more options when using nonlinear regression. Therefore, it often makes sense to use Prism's nonlinear regression analysis to fit a straight line. In particular, the nonlinear regression analysis lets you:

- Fit to both a linear and nonlinear model, and compare the two models.
• Apply differential weighting.

• Automatically exclude outliers.

• Use a robust fitting method.

• Perform a normality test on the residuals.

• Inspect the correlation matrix or dependencies.

• Compare the scatter of points from the line with the scatter among replicates with a replicates test.

• Report the best-fit values with 90% confidence limits (or any others). Prism's linear regression analysis only reports 95% CI. Nonlinear regression lets you choose the confidence level you want.

• Report the results of interpolation from the line/curve along with 95% confidence intervals of the predicted values. Prism's linear regression analysis does not include those confidence intervals.

• With linear regression, the SE of the slope is always reported with the slope as a plus minus value. With nonlinear regression, the SE values are a separate block of results that can be copy and pasted elsewhere.

• Use global nonlinear regression to fit one line to several data sets. Or share the intercept or slope among several data sets, while fitting the other parameter individually to each data set.

• Run a Monte Carlo analysis.

• When you enter data with multiple replicates at each X value, Prism's nonlinear regression can perform the replicates test to ask whether the data deviate systematically from the straight line model. Prism does not offer the replicates test with linear regression.

• Test whether the slope (or intercept) significantly differs from some proposed value. For example, test whether the slope differs from a hypothetical value of 1.0, or whether an intercept differs significantly from 0.0.
Lines that must be fit with nonlinear regression

There are two situations where you might first think that linear regression is the best analysis, but in fact nonlinear regression is necessary:

- If your Y axis uses a logarithmic or probability scale, then a straight line on the graph is created by a nonlinear model. In this case, although the line on the graph is straight, the model is not actually linear. You need to fit the 'line' with nonlinear regression.

- If you want to fit two lines to different segments of the data, this cannot be done with Prism's linear regression analysis. However, Prism's nonlinear regression can fit segmental linear regression.

3.1.1.4 Confidence and prediction bands (linear regression)

Plotting confidence or prediction bands

If you check the option box on the top of the Linear regression parameters dialog, Prism will calculate and graph either the 95% confidence band or 95% prediction band of the regression line.
To adjust the appearance of the confidence or prediction bands, go to the Format Graph dialog, select the dataset that represents the regression line, and adjust the error bars and area fill settings. You can also choose to fill the area enclosed by the confidence or prediction bands.

Plotting both confidence and prediction bands on the same graph

Prism lets you choose either a confidence band or a prediction band as part of the linear regression dialog. But not both. To plot both on one
graph, you need to analyze your data twice, choosing a confidence band the first time and a prediction band the second time.

The regression lines (and bands) are data sets that you can add to any graph by drag-n-drop or using the Change menu from the graph. So go to the graph with the first regression line, and drag the second regression analysis onto that graph. Or double click on the graph, and use the middle tab of Format Graph to add the other regression data set.

**Meaning of confidence bands**

Two confidence bands surrounding the best-fit line define the confidence interval of the best-fit line.

![Image](image1)

The dashed confidence bands are curved. This does not mean that the confidence band includes the possibility of curves as well as straight lines. Rather, the curved lines are the boundaries of all possible straight lines. The figure below shows four possible linear regression lines (solid) that lie within the confidence band (dashed).

![Image](image2)

Given the assumptions of linear regression, you can be 95% confident that the two curved confidence bands enclose the true best-fit linear
regression line, leaving a 5% chance that the true line is outside those boundaries.

Many data points will be outside the 95% confidence bands. The confidence bands are 95% sure to contain the best-fit regression line. This is not the same as saying it will contain 95% of the data points.

**Meaning of the prediction bands**

Prism can also plot the 95% prediction bands. The prediction bands are further from the best-fit line than the confidence bands, a lot further if you have many data points. The 95% prediction band is the area in which you expect 95% of all data points to fall. In contrast, the 95% confidence band is the area that has a 95% chance of containing the true regression line. This graph shows both prediction and confidence intervals (the curves defining the prediction intervals are further from the regression line).

![Graph showing regression line and confidence bands](image)

**When to plot confidence and prediction bands**

The confidence bands sort of combine the confidence intervals of the slope and intercept in a visual way. Use confidence bands to learn how precisely your data define the best-fit line.

Prediction bands are wider, to also include the scatter of the data. Use prediction bands when your main goal is show the variation in your data.
3.1.1.5 Graphing tips: Linear regression

Graphing the regression line

When Prism performs linear regression, it automatically superimposes the line on the graph.

If you need to create additional graphs, or change which line is plotted on which graph, keep in mind that the line generated by linear regression is seen by Prism as a data set. You can add lines to a graph or remove lines from a graph on the 'Data sets on graph' tab of the Format Graph dialog.

Residuals

If you check an option on the linear regression dialog, Prism will create a results table with residuals, which are the vertical distances of each point from the regression line. The X values in the residual table are identical to the X values you entered. The Y values are the residuals. A residual with a positive value means that the point is above the line; a residual with a negative value means the point is below the line.

When Prism creates the table of residuals, it also automatically makes a new graph containing the residuals and nothing else. You can treat the residuals table like any other table, and do additional analyses or make additional graphs.

If the assumptions of linear regression have been met, the residuals will be randomly scattered above and below the line at Y=0. The scatter should not vary with X. You also should not see large clusters of adjacent points that are all above or all below the Y=0 line.

See an example of residuals from nonlinear regression.

3.1.2 Results of linear regression

3.1.2.1 Slope and intercept

How Prism reports the slope and intercept

Prism first reports the best-fit values of the slope and intercept, along with their standard errors. It also reports the X intercept and the reciprocal of the slope. Below those values, it reports the 95% confidence interval of the slope and both intercepts.
At the bottom of the results page, the slope and intercept are reported again in the form of the equation that defines the best-fit line. You can copy this equation and paste onto a graph, or into a manuscript.

Interpreting the slope and intercept

The slope quantifies the steepness of the line. It equals the change in Y for each unit change in X. It is expressed in the units of the Y-axis divided by the units of the X-axis. If the slope is positive, Y increases as X increases. If the slope is negative, Y decreases as X increases.

The Y intercept is the Y value of the line when X equals zero. It defines the elevation of the line.
Interpreting the standard errors and confidence intervals

The standard error values of the slope and intercept can be hard to interpret, but their main purpose is to compute the 95% confidence intervals.

If you accept the assumptions of linear regression, there is a 95% chance that the 95% confidence interval of the slope contains the true value of the slope, and that the 95% confidence interval for the intercept contains the true value of the intercept. The width of the confidence intervals is determined by the number of data points, their distances from the line, and the spacing of the X values.

Prism can also plot either the confidence or prediction bands.

3.1.2.2 r², a measure of goodness-of-fit of linear regression

The meaning of r²

The value r² is a fraction between 0.0 and 1.0, and has no units. An r² value of 0.0 means that knowing X does not help you predict Y. There is no linear relationship between X and Y, and the best-fit line is a horizontal line going through the mean of all Y values. When r² equals 1.0, all points lie exactly on a straight line with no scatter. Knowing X lets you predict Y perfectly.

How r² is computed

This figure demonstrates how Prism computes r².
The left panel shows the best-fit linear regression line. This line minimizes the sum-of-squares of the vertical distances of the points from the line. Those vertical distances are also shown on the left panel of the figure. In this example, the sum of squares of those distances (SSreg) equals 0.86. Its units are the units of the Y-axis squared. To use this value as a measure of goodness-of-fit, you must compare it to something.

The right half of the figure shows the null hypothesis -- a horizontal line through the mean of all the Y values. Goodness-of-fit of this model (SStot) is also calculated as the sum of squares of the vertical distances of the points from the line, 4.907 in this example. The ratio of the two sum-of-squares values compares the regression model with the null hypothesis model. The equation to compute $r^2$ is shown in the figure. In this example $r^2$ is 0.8428. The regression model fits the data much better than the null hypothesis, so SSreg is much smaller than SStot, and $r^2$ is near 1.0. If the regression model were not much better than the null hypothesis, $r^2$ would be near zero.

You can think of $r^2$ as the fraction of the total variance of Y that is "explained" by variation in X. The value of $r^2$ (unlike the regression line itself) would be the same if X and Y were swapped. So $r^2$ is also the fraction of the variance in X that is "explained" by variation in Y. In other words, $r^2$ is the fraction of the variation that is shared between X and Y.

In this example, 84% of the total variance in Y is "explained" by the linear regression model. The variance (SS) of the data from the linear regression model equals only 16% of the total variance of the Y values (SStot).

**Why Prism doesn't report $r^2$ in constrained linear regression**

Prism does not report $r^2$ when you force the line through the origin (or any other point), because the calculations would be ambiguous. There are two ways to compute $r^2$ when the regression line is constrained. As you saw in the previous section, $r^2$ is computed by comparing the sum-of-squares from the regression line with the sum-of-squares from a model defined by the null hypothesis. With constrained regression, there are two possible null hypotheses. One is a horizontal line through the mean of all Y values. But this line doesn't follow the constraint -- it does
not go through the origin. The other null hypothesis would be a horizontal line through the origin, far from most of the data.

Because \( r^2 \) is ambiguous in constrained linear regression, Prism doesn't report it. If you really want to know a value for \( r^2 \), use nonlinear regression to fit your data to the equation \( Y = \text{slope} \times X \). Prism will report \( r^2 \) defined the first way (comparing regression sum-of-squares to the sum-of-squares from a horizontal line at the mean \( Y \) value).

**Upper or lower case?**

With linear regression, it is conventional to use the abbreviation \( r^2 \). With nonlinear regression, the convention is to use \( R^2 \). There appears to be no reason for this distinction.

**3.1.2.3 Is the slope significantly different than zero?**

**Interpreting the P value when Prism fits both slope and intercept**

Prism reports the P value testing the null hypothesis that the overall slope is zero. The P value answers this question:

If there were no linear relationship between \( X \) and \( Y \) overall, what is the probability that randomly selected points would result in a regression line as far from horizontal (or further) than you observed?

Equivalently:

If there were no linear relationship between \( X \) and \( Y \) overall, what is the probability that randomly selected points would result in an \( R^2 \) value as high (or further) as you observed?

The P value is calculated from an F test, and Prism also reports the value of F and its degrees of freedom. You would get exactly the same P value from the t ratio computed by dividing the slope by its standard error.

**Interpreting the P value when Prism fits only the slope**

Beware of the P value from linear regression when you force the line to go through the origin or some other point. It is easy to misinterpret. The graph below demonstrates the problem:
The data points almost form a horizontal line.

When fit with linear regression the usual way (fit both slope and intercept; green line), the best fit value of the slope is 0.00. The P value answers the question: If the true slope is zero, what is the chance that the slope will be further from zero than the observed slope due only to random sampling. Since the observed slope is zero, there is almost a 100% chance of obtaining a slope that is further than zero than observed! So the P value is greater than 0.99, as high as a P value can be. Some people are confused and think the P value should be small purely because the points for a pattern. Not so. The P value, from conventional linear regression fitting both slope and intercept, will be small only when the points form a linear pattern that is not horizontal.

The results are very different when you fit linear regression with the constraint that the line has to go through the origin (blue line). To make the line go through the origin and also go near the points, the best-fit line has a slope far from zero. Since the line is far from horizontal, the P value is tiny. Given the constraint that the line must go through the origin (X=0, Y=0; lower-left of graph), the data are quite convincing that best fit line is far from horizontal, so it makes sense that the P value is tiny.

Constraining a line to go through the origin (or some other point) can be very useful in some situations. Usually this option is used to fit calibration curves used for interpolation, in which case the P value is not useful. If you force the line through the origin, be very wary when interpreting the P value. It is rarely useful, and easy to misinterpret.

Download the Prism file.
3.1.2.4 Comparing slopes and intercepts

Prism compares slopes of two or more regression lines if you check the option: "Test whether the slopes and intercepts are significantly different".

**Comparing slopes**

Prism compares slopes first. It calculates a P value (two-tailed) testing the null hypothesis that the slopes are all identical (the lines are parallel). The P value answers this question:

If the slopes really were identical, what is the chance that randomly selected data points would have slopes as different (or more different) than you observed.

**If the P value is less than 0.05**

If the P value is low, Prism concludes that the lines are significantly different. In that case, there is no point in comparing the intercepts. The intersection point of two lines is:

\[
X = \frac{\text{Intercept}_1 - \text{Intercept}_2}{\text{Slope}_2 - \text{Slope}_1}
\]

\[
Y = \text{Intercept}_1 + \text{Slope}_1 \cdot X = \text{Intercept}_2 + \text{Slope}_2 \cdot X
\]

**If the P value for comparing slopes is greater than 0.05**

If the P value is high, Prism concludes that the slopes are not significantly different and calculates a single slope for all the lines. Essentially, it shares the Slope parameter between the two data sets.

**Comparing intercepts**

If the slopes are significantly different, there is no point comparing intercepts. If the slopes are indistinguishable, the lines could be parallel with distinct intercepts. Or the lines could be identical with the same slopes and intercepts.

Prism calculates a second P value testing the null hypothesis that the lines are identical. If this P value is low, conclude that the lines are not identical (they are distinct but parallel). If this second P value is high, there is no compelling evidence that the lines are different.
**Relationship to ANCOVA and global regression**

This method is equivalent to an Analysis of Covariance (ANCOVA), although ANCOVA can be extended to more complicated situations. It also is equivalent to using Prism's nonlinear regression analysis with a straight-line model, and using an F test to compare a global model where slope is shared among the data sets with a model where each dataset gets its own slope.

**Reference**


**3.1.2.5 Runs test following linear regression**

The runs test determines whether your data differ significantly from a straight line. Prism can only calculate the runs test if you entered the X values in order.

A run is a series of consecutive points that are either all above or all below the regression line. In other words, a run is a consecutive series of points whose residuals are either all positive or all negative.

If the data points are randomly distributed above and below the regression line, it is possible to calculate the expected number of runs. If there are Na points above the curve and Nb points below the curve, the number of runs you expect to see equals \([(2NaNb)/(Na+Nb)]+1\). If you observe fewer runs than expected, it may be a coincidence of random sampling or it may mean that your data deviate systematically from a straight line. The P value from the runs test answers this question:

If the data really follow a straight line, and you performed many experiments like this one, what fraction of the time would you obtain as few (or fewer) runs as observed in this experiment?

If the runs test reports a low P value, conclude that the data do not really follow a straight line, and consider using nonlinear regression to fit a curve.
The P values are always one-tail, asking about the probability of observing as few runs (or fewer) than observed. If you observe more runs than expected, the P value will be higher than 0.50.

3.1.2.6 Analysis checklist: Linear regression

- **Can the relationship between X and Y be graphed as a straight line?**

  In many experiments the relationship between X and Y is curved, making linear regression inappropriate. It rarely helps to transform the data to force the relationship to be linear. Better, use nonlinear curve fitting.

- **Is the scatter of data around the line Gaussian (at least approximately)?**

  Linear regression analysis assumes that the scatter of data around the best-fit line is Gaussian.

- **Is the variability the same everywhere?**

  Linear regression assumes that scatter of points around the best-fit line has the same standard deviation all along the curve. The assumption is violated if the points with high or low X values tend to be further from the best-fit line. The assumption that the standard deviation is the same everywhere is termed homoscedasticity. (If the scatter goes up as Y goes up, you need to perform a weighted regression. Prism can't do this via the linear regression analysis. Instead, use nonlinear regression but choose to fit to a straight-line model.)

- **Do you know the X values precisely?**

  The linear regression model assumes that X values are exactly correct, and that experimental error or biological variability only affects the Y values. This is rarely the case, but it is sufficient to assume that any imprecision in measuring X is very small compared to the variability in Y.

- **Are the data points independent?**

  Whether one point is above or below the line is a matter of chance, and does not influence whether another point is above or below the line.
**Are the X and Y values intertwined?**

If the value of X is used to calculate Y (or the value of Y is used to calculate X) then linear regression calculations are invalid. One example is a Scatchard plot, where the Y value (bound/free) is calculated from the X value. Another example would be a graph of midterm exam scores (X) vs. total course grades (Y). Since the midterm exam score is a component of the total course grade, linear regression is not valid for these data.

### 3.1.2.7 Questions and answers

- **What is the difference between correlation and linear regression?**
  
  *The answer is in the Correlation section.*

- **Why is the 95% confidence interval of the X intercept asymmetrical?**

  The 95% confidence interval for the X-intercept is not symmetrical around the X-intercept. It goes further in one direction than the other, as illustrated in the graph below.

  ![Graph showing asymmetrical 95% confidence interval](image)

  Follow the Y=0 baseline from left to right. The region between the 95% confidence bands for the best fit line (blue curves) is the 95% CI of the X intercept. You can see that this confidence interval (between the two
outmost dotted lines) is not symmetrical around the X intercept (the middle dotted line).

This asymmetry will be very noticeable if you only have a few points with lots of scatter, and will be almost unnoticeable with lots of points with little scatter.

GraphPad Prism reports the 95% confidence interval of the X intercept if you check an option on the Linear regression parameters dialog.

Because the uncertainty is not symmetrical, it rarely makes sense to report a standard error of the X-intercept. It is much better to report both ends of the 95% confidence interval, which Prism reports. If you really want to compute a single standard error for the X intercept, you can do so by choosing nonlinear regression, and fitting this user-defined equation to the data:

\[ Y = \text{slope}(X - \text{Xintercept}) \]

Prism will report the best-fit value of the X intercept along with a SE and 95% confidence interval. Since this confidence interval will be computed from the SE value it will be symmetrical around the X intercept, and so won't be as accurate as the asymmetrical interval reported by linear regression.

How to test whether the slope from linear regression differs from 1.0 or some other value?

Prism tests whether the best-fit slope from linear regression line differs significantly from zero. But you need to use extra steps to test whether the slope differs from some other value.

Since Prism's linear regression analysis can't answer the question, use nonlinear regression instead. Nonlinear regression can fit a straight line and offers many advantages.

In the nonlinear regression parameters dialog, go to the Compare tab, check the option to ask compare the best-fit value of a parameter to a theoretical value, and then choose Slope and enter the theoretical value (1.0).

Can Prism do weighted linear regression?

Prism's linear regression analysis cannot handle differential weighting. However, Prism's nonlinear regression analysis can do weighted
regression, and you can fit a straight line using "nonlinear" regression.

How can Prism fit linear regression with the slope constrained to equal 1.0 (or some other value)?

Prism's linear regression analysis cannot constrain the slope to a particular value. However, Prism's nonlinear regression analysis can constrain any parameter to any value, and you can fit a straight line using "nonlinear" regression.

How to fit one linear regression line to multiple data sets?

Prism usually fits one line through each data set. But you can get it to fit one line through all the data. Use the nonlinear regression analysis to fit the line. Then go to the Constrain tab, and share both the slope and the intercept. "Sharing" means that Prism fits one best-fit value for all the data sets, rather than one for each data set. Since you share all the parameters, only one line is fit to all the data.

Why doesn't Prism report R2 for linear regression when I force the line through the origin (or some other point)?

The problem is that when you constrain a line to go through a point, there would be two ways to compute R²:

- Compare the fit of the best-fit line with the fit of a horizontal line at the mean Y value. But that null hypothesis (horizontal line through the Ymean) doesn't obey the constraint that it go through the origin.

- Compare the best-fit line with a horizontal line at Y=0. This obeys the constraint, but often fits the data really badly, pushing up the R² value.

Since R² is ambiguous when you constrain linear regression, Prism simply does not report it when you force the line through the origin (or other point).

If you fit the line with nonlinear regression, Prism will report R² using the first definition above.

Fitting the crossing point of two intersecting linear regression lines.
Enter your data onto one data table, with Y values in columns A and B next to corresponding X values in the X column. If the X values don't match for the two data sets, just leave some Y values blank.

Use nonlinear regression, and enter this user-defined equation.

\[ Y = Y_{cross} + (X - X_{cross}) \times \text{Slope} \]

This equation defines the Y coordinate of any point along either line to equal the Y coordinate of the crossing point plus the slope times the X distance from the crossing point to that point.

Set these rules for initial values:

- \( Y_{cross} \): 1*Ymid (the average of the highest and lowest Y values)
- \( X_{cross} \): 1*Xmid
- \( \text{Slope} \): 1*(Ymax-Ymin)/(Xmax-Xmin).

Define \( X_{cross} \) and \( Y_{cross} \) to both be shared parameters, so Prism will find one global best-fit value for those parameters that applies to both data sets. Do not share the slope parameter, as you want Prism to fit separate slopes for each data set. Getting these sharing settings correct is essential.

Prism will fit four parameters:

- Slope for the first data set
- Slope for the second data set
- \( X_{cross} \) (shared for both data sets)
- \( Y_{cross} \) (shared for both data sets).

That makes sense. You have two lines, and it takes two parameters (usually a slope and intercept) to describe each. Here, the equations were rearranged so the program doesn't fit two separate Y intercepts, but rather fits the X and Y values of the crossing point.

Here is an example, with the key results circled. Prism file.
3.1.3 Deming regression

3.1.3.1 Key concepts: Deming regression

Standard linear regression assumes that you know the X values perfectly, and all the uncertainty is in Y. It minimizes the sum of squares of the vertical distance of the points from the line.

If both X and Y variables are subject to error, fit linear regression using a method known as Deming, or Model II, regression.

If your goal is to compare two analysis methods, consider using a Bland-Altman plot instead.
3.1.3.2 How to: Deming regression

1. Create a data table

From the Welcome or New Table dialog, choose to create XY data table.

If you are just getting started, choose the sample data: Linear regression -- Compare slopes.

If you are entering your own data, choose a to enter a single Y value for each point (no replicates, no error values).

2. Enter data

If you enter Y values for several data sets (column A, B and C), Prism will report regression results for X vs. YA, for X vs. YB, and for X vs. YC.

3. Analysis choices

Click Analyze and choose Deming regression from the list of XY analyses.

Most of the dialog is self-explanatory. The choices on top are the hardest to understand, but also are the most important as the calculations depend on knowing relative magnitude of the errors in X and Y.

X and Y have equal uncertainty

If you are comparing two analytical methods, X and Y, assessed in the same units, you will probably find it reasonable to assume that both share the same standard deviation. In this case, check the first option. There is no need to estimate that standard deviation. It is enough to declare that X and Y have equal standard deviations. In this case, Deming regression minimizes the sum of the squares of the
perpendicular distances of the points from the line. This is also called \textit{orthogonal regression}.

\textbf{X and Y have different uncertainties}

If you are not willing to assume that X and Y have the same amount of uncertainty, then you need to enter the SD of each. Enter the SD of X values in the same units as X values are entered, and the SD of Y values in the same units as Y values are entered. Don't enter the SD of all X (or Y) values you enter. Enter the average SD (determined separately) of repeated measurements of X and Y.

How do you know what values to enter? To assess the uncertainty (error) of a method, collect duplicate measurements from a number of samples using that method. Calculate the standard deviation of the error using the equation below (1), where each \( d_i \) is the difference between two measurements of the same sample (or subject), and \( N \) is the number of measurements you made (\( N \) equals twice the number of samples, since each sample is measured twice).

\[
SD_{\text{error}} = \sqrt{\frac{\sum d_i^2}{N}}
\]

Compute the SD of the error for each method or variable (X and Y), enter the two \( SD_{\text{error}} \) values into the Deming regression analysis dialog, and Prism will fit the line for you. If the X variable has a much smaller SD than the Y value, the results will be almost identical to standard linear regression.

If you try to compare Prism’s results with those of another program or book, you may encounter the variable \( \lambda \) (lambda), which quantifies the inequality between X and Y errors.

\[
\lambda = \left( \frac{SD_{X\text{error}}}{SD_{Y\text{error}}} \right)^2
\]

Prism requires you to enter individual SD values, but uses these values only to calculate \( \lambda \), which is then used in the Deming regression calculations. If you know \( \lambda \), but not the individual SD values, enter the square root of \( \lambda \) as the SD of the X values, and enter 1.0 as the SD of the Y error. The calculations will be correct, since Prism uses those two values only to compute \( \lambda \).
3.1.3.3 Q&A: Deming Regression

Why doesn't Prism report $R^2$ with Deming Regression results?

When Prism performs Deming regression, it reports the slope and intercepts with confidence intervals, and reports a P value testing the null hypothesis that the slope is really zero. However, Prism does not report any measure of goodness-of-fit with Deming regression, and so does not report $R^2$. The reason is that we have been unable to find any paper or text that would explain how to compute or interpret such a value. In ordinary linear or nonlinear regression, $R^2$ is the fraction of the variation that is accounted for by the model. But with Deming regression, this definition doesn't really make sense, and it isn't obvious to us how to extend it.

Is Deming regression the same as orthogonal linear regression?

In Prism's Deming dialog, you specify whether X and Y are in the same units with equal uncertainties (variation). If you choose this option, Deming regression minimizes the sum of the square of the perpendicular distances of the points from the line. This is also called *orthogonal linear regression*. If you specify different SD values for X and Y, then Deming regression is not the same as orthogonal linear regression.

How was the equation to compute the SD from a set of duplicate measurements derived?

The previous page showed the equation for assessing the uncertainty (error) of a method from duplicate measurements from a number of samples using that method:

$$SD_{\text{error}} = \sqrt{\frac{\sum d_i^2}{N}}$$

It is easy to derive that equation. To compute the variance, you find the average square of the difference between each value and the mean. For pairs, the mean is half the difference. So take half that distance squared, but also compute it for the other value of the pair. For both values, therefore, the contribution is $d_i$ squared. Add that up for all the
measurements and divide by the number of measurements, and you have the variance. Take the square root to get the SD.

Why N, rather than N-1? I suspect it should be N-1, but that won't matter much. It will make the SD a bit smaller than it should be, but that will be true for both the X and Y SD, so the ratio will hardly be changed.

3.1.3.4 Analysis checklist: Deming regression

✔ Can the relationship between X and Y be graphed as a straight line?

In many experiments the relationship between X and Y is curved, making linear regression inappropriate. It rarely helps to transform the data to force the relationship to be linear. Better, use nonlinear curve fitting.

✔ Are the data points independent?

Whether one point is above or below the line is a matter of chance, and does not influence whether another point is above or below the line.

✔ Are the X and Y values intertwined?

If the value of X is used to calculate Y (or the value of Y is used to calculate X) then linear regression calculations are invalid. One example is a Scatchard plot, where the Y value (bound/free) is calculated in part from the X value (bound). Another example would be a graph of midterm exam scores (X) vs. total course grades (Y), since Y is in part computed from X.

✔ Do you know the relative uncertainty of X and Y?

Ordinary linear regression assumes that you know the X values perfectly, and all the uncertainty is in Y. Deming regression assumes there is uncertainty in both variables. You have to specify the relative uncertainties, either by specifying that X and Y are equally uncertain or by entering the SD of each. If these values are incorrect, the Deming results won't be useful.

3.2 Interpolating from a standard curve

Prism makes it very easy to interpolate unknown values from a standard curve. Enter the standards with both X and Y values, fit a line or curve, and Prism will tell you which X values correspond to Y values you entered on the same data table.
3.2.1  **Key concept: Interpolating**

A common use of nonlinear (and linear) regression is interpolating. The basic idea is simple:

1. Enter a standard curve. Each point consists of a concentration (or logarithm of concentration) and a response. The responses are commonly radioactivity, fluorescence, or optical density.

2. Fit a line or curve through the standards. For this purpose, you don’t much care about the meaning of the model, or the interpretation of the best-fit values of the parameters (EC50, etc.).

3. For unknown samples, you know the response, but not the concentration. So read the concentration from the curve:
Notes:

- This graph shows the response going up as the concentration gets larger. In many cases, the response goes down as the concentration gets larger. Interpolation works in either case.

- Sigmoid shaped dose-response curves are commonly used with many assays. If the X values analyzed by the analysis are the logarithm of concentration, then so are the interpolated values. You need to take the antilogarithm of these values to return to concentration units.

- This example shows the most common use of interpolating. Finding an X value (concentration) for a given Y value. Prism can also interpolate the other way: Finding a Y value for an entered X value.

3.2.2 How to interpolate

Prism is not designed to analyze any particular kind of assay

Prism is a very general program, used by scientists in many fields. It is easy for you to interpolate standard curves using Prism, but you must make some decisions. Unlike some specialized programs, Prism is not set up so you can blindly enter data and get results without thinking.
Entering data for interpolation

Enter the standard curve in the top rows of a table, and then enter the unknown values right below those standards. In most cases, as shown below, you'll enter Y values and leave X blank. Prism can also interpolate the other way. In this case, enter X values and leave Y blank.

<table>
<thead>
<tr>
<th>Table format</th>
<th>X</th>
<th>A:Y1</th>
<th>A:Y2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>log</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>8</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>9</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>11</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Entering labels for the unknowns, as shown below, is optional.

Note that it is essential that the unknowns go below the standard curve values. If you place the unknowns above the standard curve, Prism will not interpolate. Of course, the unknown Y values must be in the same units as the Y values you entered for the standard curve.

Fitting a standard curve and interpolating

Four analyses in Prism let you interpolate values from curves. The top row of buttons in the analysis block provide shortcuts to linear regression, nonlinear regression, and the simplified analysis for interpolating a standard curve.

Interpolate a standard curve

This analysis (new to Prism 6) is a simple way to interpolate using nonlinear regression.
The choices on the dialog are simple. Choose a model to fit to your data. The most commonly used models for standard curves are shown in the dialog. Next choose how to handle outliers. You can choose to use robust regression so outliers have little impact. Or you can ask Prism to identify, and perhaps eliminate, outliers using the ROUT method. Finally, if the scatter among replicates is larger when the Y values are larger, consider using relative weighting.

Note that this simplified analysis to interpolate is simply a subset of the nonlinear regression analysis, with fewer choices on the dialog. If you want to select options that aren't available, click the "More..." button at the bottom of the dialog to convert the analysis to the full nonlinear regression analysis, with many more options (six tabs worth). Once you switch to nonlinear regression analysis, it is not possible to switch back (except by starting over).

**Nonlinear regression**

Nonlinear regression offers the interpolation option at the bottom of the first (Fit) tab of the dialog. This is a good choice if you want to use options not available on the simpler interpolation analysis.

**Linear regression**

Linear regression offers a checkbox to interpolate the top of the dialog.

**Spline**

Prism can also interpolate from a standard curve in the spline analysis. Splines go through every point. The advantage of a spline is that you don't have to choose a model to fit to your data. The disadvantage is that the curve may wiggle too much.

### 3.2.3 How to graph the interpolated values

The table that includes the interpolated values has a greed grid which means it can be used to make graphs. While looking at that table, drop New... Graph of existing data. Choose an XY graph and you are done.

### 3.2.4 Example: Interpolating from a sigmoidal standard curve

**1. Create the data table**
From the Welcome or New Table dialog, choose to create an XY data table, and select the sample data set: RIA or ELISA. Interpolate unknowns from sigmoidal curve.

2. Inspect the data

The sample data may be partly covered by a floating note explaining how to fit the data (for people who are not reading this help page). You can move the floating note out of the way, or minimize it.

The first seven rows contain the standard curve, in duplicate. Below that are three unknown values. These have a Y values that you measured, but no X. The goal of this analysis is to interpolate the corresponding X values (concentrations) for these unknowns. Note that three of the four unknowns are labeled, so you can later match up the results with the labels.

Why are X values negative? Because in this example, the X values are the logarithm of concentrations expressed in molar. So a concentration of 1 micromolar ($10^{-6}$ Molar) would be entered as -6.
3. View the graph

The graph Prism makes automatically is fairly complete. You can customize the symbols, colors, axis labels, etc. You can also choose to plot the individual duplicates rather than plot the means. Since the unknowns have no X value, they are not included on the graph.

4. Choose the standard curve analysis

Click the Analyze button and from the list of XY analyses choose: Interpolate a Standard Curve.

Alternatively, you can click the “Interpolate a standard curve” button right on top of the Analyze button.

5. Choose a model

Choose the equation: Sigmoidal, 4PL, X is log(Concentration). Note that 4PL means four parameter logistic, which is another name for this kind of equation.
For this example, leave all the other settings to their default values.

Click OK to see the curves superimposed on the graph.

6. Inspect the results

The results appear on several pages.

The first page shows you the interpolated values.
The second page is the table of results for the overall curve fit. It tabulates the best-fit values of the parameters and much more. For this example, we aren't too interested in these results.

7. Transform the results

The X column of the results table has the interpolated values we want. These are in the same units as the X values, so are the logarithm of concentration. Prism can transform these values to concentration units.

Click the Analyze button and choose Transform at the top of the Analyze dialog.

On the Transform dialog check the option to transform X values and choose the transform $X=10^X$

Now the X column is in molar concentration units. Note that the column title hasn't changed. Prism isn't smart enough to adjust the column titles when you transform data.

Click and edit the column title to "Concentration (M)".

3.2.5 Equations used for interpolating

You can interpolate from any curve fit through a set of standards. However, the equations listed below are used most commonly. These equations are instantly available from the interpolation analysis. You'll
also find this set of equations in the equation tree used in nonlinear regression.

Notes:

- These graphs show the curve going up (increasing Y) as X increases. But all will work for downward sloping curves too.

- All of these equations (except Pade) are also present in other equation folders in the nonlinear regression dialog.

**Sigmoidal, 4PL**,  
X is log(concentration)

Note that in nonlinear regression this equation is called "log(agonist) vs. response -- Variable slope"

**Asymmetric Sigmoidal, 5PL**,  
X is log(concentration)

Note that in nonlinear regression this equation is called "log(agonist) vs:"

**Biphasic**

Note that in nonlinear regression this equation is called "Asymmetrical"
**Straight line**

\[ Y = B_0 + B_1 X + B_2 X^2 \]

**Semilog line**

- X is log, Y is linear

**Hyperbola**

- X is concentration

**Padé (1,1) approximant**

**Second order polynomial (quadratic)**

\[ Y = B_0 + B_1 X + B_2 X^2 \]
3.2.6 The results of interpolation

Where are the interpolated results?

Prism places the interpolation results into its an additional results page (or two additional pages).

<table>
<thead>
<tr>
<th>X</th>
<th>RIA data log(Concentration) (Interpolated)</th>
<th>RIA data Data Set A (Entered)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Unknown 1</td>
<td>-6.549</td>
<td>1123.000</td>
</tr>
<tr>
<td>2 Unknown 2</td>
<td>-7.175</td>
<td>1345.000</td>
</tr>
<tr>
<td>3 Unknown 3</td>
<td>-7.573</td>
<td>1456.000</td>
</tr>
<tr>
<td>4</td>
<td>-6.408</td>
<td>987.000</td>
</tr>
</tbody>
</table>

To view this table, click the interpolated results page of the results. In some cases you may first need to click the + in front of the results main page to see all the pages within.
What units are the results in?

The interpolated results are expressed in the same units as the standard curves. If you entered a Y value, and Prism interpolated an X value, then that X value is in the same units as the standards you entered.

If the X values in the table are the logarithm of concentration (common), then the interpolated results are also logarithms of concentration.

In some assays, different samples are diluted differently. Prism does not correct for dilution, so you'll need to make those corrections manually.

3.2.7 Interpolating with replicates in side-by-side subcolumns

Choices when entering data with replicates in subcolumns

It is common to enter replicate values for the unknowns, as shown below.

<table>
<thead>
<tr>
<th>Table format</th>
<th>X (log Concentration)</th>
<th>Group A</th>
<th>Data Set-A</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>X</td>
<td>A1:Y1</td>
<td>A1:Y2</td>
</tr>
<tr>
<td>1</td>
<td>Title</td>
<td>1597</td>
<td>1531</td>
</tr>
<tr>
<td>2</td>
<td>Title</td>
<td>1453</td>
<td>1471</td>
</tr>
<tr>
<td>3</td>
<td>Title</td>
<td>1314</td>
<td>1246</td>
</tr>
<tr>
<td>4</td>
<td>Title</td>
<td>751</td>
<td>771</td>
</tr>
<tr>
<td>5</td>
<td>Title</td>
<td>336</td>
<td>366</td>
</tr>
<tr>
<td>6</td>
<td>Title</td>
<td>328</td>
<td>212</td>
</tr>
<tr>
<td>7</td>
<td>Title</td>
<td>207</td>
<td>367</td>
</tr>
<tr>
<td>8</td>
<td>Unknown 1</td>
<td>1123</td>
<td>1211</td>
</tr>
<tr>
<td>9</td>
<td>Unknown 2</td>
<td>1345</td>
<td>1298</td>
</tr>
<tr>
<td>10</td>
<td>Unknown 3</td>
<td>1456</td>
<td>1523</td>
</tr>
<tr>
<td>11</td>
<td>Title</td>
<td>987</td>
<td>1111</td>
</tr>
</tbody>
</table>

Since Prism's results tables cannot be configured to put side-by-side replicates in the X column, you have two choices for how to organize the results. These choices are available at the bottom of the Output tab of the nonlinear regression dialog.

Location of interpolated X values

- X column, with replicate values stacked
- Y column, maintaining the side-by-side arrangement of replicates
Place computed unknown values into the X column, with replicate values stacked

In this case, Prism creates two pages of results.

The interpolation is the same for both tables. Prism interpolates each replicate value individually.

The first page, Interpolated Mean X Values, shows the mean of the Y values you entered and the mean of the interpolated X values. The X column shows the concentrations (or logarithm of concentrations) to match the data you entered.

<table>
<thead>
<tr>
<th>X</th>
<th>A</th>
</tr>
</thead>
<tbody>
<tr>
<td>log(Concentration): (Interpolated)</td>
<td>RIA data: Set-A. (Entered)</td>
</tr>
<tr>
<td>1 Unknown 1</td>
<td>-6.735</td>
</tr>
<tr>
<td>2 Unknown 2</td>
<td>-7.103</td>
</tr>
<tr>
<td>3 Unknown 3</td>
<td>-7.554</td>
</tr>
<tr>
<td>4</td>
<td>-6.516</td>
</tr>
</tbody>
</table>

The second page, Interpolated X replicates, shows individual replicate values for the interpolated results. The X column maintains the same meaning as it does in the data table you entered. To do so, with only a single X column, requires that the replicates be stacked.
Place computed unknown values into a Y data set, maintaining the side-by-side arrangement of replicates.

When you make this choice, the interpolated values appear in the Y column, and there is no X column. You can view the interpolated values, with their name (if you entered row titles). But the Y values you entered are not shown.

The first page shows the mean values. The second page shows the interpolated values in side-by-side replicates, to match the format you entered them. Even though these values are interpolated X values, note that the results are in the Y column.

3.2.8 Interpolating several data sets at once

Prism can interpolate from several data sets at once. To do this, put all the unknowns on lower rows on the data table than all the knowns (left panel). Prism won’t be able to interpolate past the first data set if you put the second data set below the first set of knowns (right panel).

Yes! All unknowns at the bottom.  No! Unknowns everywhere.
Why can't Prism handle the kind of data shown in the right panel above? Because then it might get mixed up about which values are unknowns to be interpolated, and which are simply adjacent to values left blank because they are missing.

### 3.2.9 When X values are logarithms

In many situations, you'll fit the standard curve to a log(dose) vs. response curve. In these cases, you'll enter the X values for the standard curve as the logarithm of concentration or you'll transform the data to logarithms before running nonlinear regression.

The interpolated X values are in the same units as the X values the analysis sees. If you are analyzing data entered or transformed so X is
the logarithm of concentration, then the interpolated values will also be logarithms.

If you want to transform these results back to concentrations, that is easy to do:

1. From the interpolated results page of results, click Analyze.
2. Choose Transform at the very top of the Analyze dialog.
3. Choose the standard list of transforms.
4. Depending on whether your interpolated values are in the X or Y column, choose to transform that column.
5. From the list of transforms, choose $Y = 10^Y$ or $X = 10^X$. These choices are about one third of the way down the long list of transforms.
6. The results sheets will have the interpolated values as concentrations, rather than logs.

Why $10^Y$ or $10^X$? Those are abbreviations for $10^Y$ or $10^X$, which are the antilogarithm functions. Review logarithms and antilogarithms.

### 3.2.10 Analysis checklist: Interpolating

Your approach in evaluating nonlinear regression depends on your goal.

In many cases, your goal is to learn from the best-fit values. If that is your goal, view a different checklist.

If your goal is to create a standard curve from which to interpolate unknown values, your approach depends on whether this is a new or established assay.

**Established assay**

If the assay is well established, then you know you are fitting the right model and know what kind of results to expect. In this case, evaluating a fit is pretty easy.

✓ *Does the curve go near the points?*
✓ Is the R² 'too low' compared to prior runs of this assay?

If so, look for outliers, or use Prism's automatic outlier detection.

✓ Are the confidence bands too wide?

The confidence bands let you see how accurate interpolations will be, so we suggest always plotting prediction bands when your goal is to interpolate from the curve. If you are running an established assay, you know how wide you expect the prediction bands to be.

**New assay**

With a new assay, you also have to wonder about whether you picked an appropriate model.

✓ Does the curve go near the points?

Look at the graph. Does it look like the curve goes near the points.

✓ Are the confidence bands too wide?

How wide is too wide? The prediction bands show you how precise interpolations will be. Draw a horizontal line somewhere along the curve, and look at the two places where that line intercepts the confidence bands. This will be the confidence interval for the interpolation.

✓ Does the scatter of points around the best-fit curve follow a Gaussian distribution?

Least squares regression is based on the assumption that the scatter of points around the curve follows a Gaussian distribution. Prism offers three normality tests (in the Diagnostics tab) that can test this assumption (we recommend the D'Agostino test). If the P value for a normality test is low, you conclude that the scatter is not Gaussian.

✓ Could outliers be impacting your results?

Nonlinear regression is based on the assumption that the scatter of data around the ideal curve follows a Gaussian distribution. This assumption leads to the goal of minimizing the sum of squares of distances of the
curve from the points. The presence of one or a few outliers (points much further from the curve than the rest) can overwhelm the least-squares calculations and lead to misleading results.

You can spot outliers by examining a graph (so long as you plot individual replicates, and not mean and error bar). But outliers can also be detected automatically. GraphPad has developed a new method for identifying outliers we call the ROUT method. You can either ask Prism to simply identify outliers or to eliminate them. These choices are adjacent on the Interpolate a Standard Curve dialog. But they are in separate tabs in the Nonlinear Regression dialog. The option to count outliers is on the Diagnostics tab, and the option to exclude outliers is on the Fit tab.

✓ Does the curve deviate systematically from the data?

If either the runs test or the replicates test yields a low P value, then you can conclude that the curve doesn't really describe the data very well. You may have picked the wrong model.

3.2.11 Reasons for blank (missing) interpolated results

No interpolated values

If the nonlinear regression procedure did not converge on a best-fit curve, then Prism will leave all interpolated values blank. It doesn't make sense to interpolate unless Prism finds a good curve.

Some interpolated values are blank

When you ask Prism to interpolate X values from Y values you enter, it will leave the result blank when it is unable to find a result. This can happen when:

• The input value for Y falls out of the vertical range of the standard curve. This happens when you fit a sigmoidal log(dose) vs response curve. If you enter an unpaired Y value that is larger than the best-fit value of Top, or smaller than the best-fit value of Bottom, then simply is no corresponding X value, and Prism will simply leave that part of the Results sheet blank. Note that "top" and "bottom" apply to the best-fit values for those fitted parameters, not the maximum and minimum
values of the input data. It simply is impossible to interpolate a X value if the Y value is too high, or too low, to be on the curve.

- The interpolated X value falls way outside the output range of X for the fitted curve. Prism will try to extrapolate a bit beyond the range of X values for which it calculates and draws the curve. But it won't extrapolate far from the ends of the curve. You can extend this range. From the Nonlinear regression Parameters dialog, choose the Range tab and enter a lower value for the minimum X and/or a higher value for the maximum X.

- You entered the unknowns in rows of the data table above (smaller row numbers) than the standard curve values. The unknowns must be entered in rows below the standards.

- You mistakenly entered the unknowns in the wrong units. The unknown values must be in the same units as the standards. If you enter unknown Y values and expect Prism to interpolate an X value, then the unknown Y values must be entered in the same units as the Y values for the standards.

**Confidence limit of interpolated value is blank**

Prism will leave the confidence interval of the interpolated value blank in two situations:

- The fit is ambiguous.

- If you chose a robust fit.

In some cases, Prism will report confidence limits for some interpolated values, but not all of them.

Prism can only report the confidence interval for an interpolated value when the curve is drawn far enough along the X axis. The graph below shows an example where Prism can report the lower confidence limit but not the upper one. The horizontal green line shows the interpolation. You asked Prism for the X value when Y=50. The interpolation works fine as does the lower confidence limit. But notice that the lower confidence band (in red) does not extend far enough to intersect the horizontal green line. That is why Prism can't report the upper confidence limit which would be the X coordinate where the lower confidence band of the curve intersects the horizontal line at Y=50. To fix this problem, go to
In other cases, the bottom confidence band bends downwards, so the upper confidence limit of the interpolated value is simply undefined. The figure below shows one such example. It doesn't matter how far you extend the X axis, the upper confidence limit of the interpolated value is simply not defined by the data. The dotted horizontal line at Y=50 will never intersect the lower of the confidence bands. The only fix would be to collect data at higher X concentrations, or perhaps more precise Y values.
3.2.12 Q&A: Interpolating

Can Prism interpolate several data sets at once?

Yes, Prism can interpolate X values from Y values entered into several data sets. Note the following:

- Prism has to figure out which values are unknowns. It uses this rule: X(or Y) values are considered as unpaired (and are used for calculation of unknowns) when they are lower in the table than all XY pairs in the data table. It is not enough that the unpaired Y values (from which you want Prism to interpolate X) is lower on the table than the paired XY values for that particular data set. They must be on lower rows in the table than all the paired XY values of all data sets in the table.

- Prism offers an option in the Output tab of nonlinear regression that let you choose where the interpolated X values should go (into
the X column staggered, or onto Y columns). If you have replicates and have unknowns in more than one data set, choose the option to show the unknowns in Y columns.

- **What units are the interpolated values expressed in?**

  When you enter a Y value, and ask Prism to interpolate the X value, that X value is in the same units as the X values you entered. Usually this is concentration. If it is the logarithm of concentration, you can transform it back to a concentration scale.

- **What happens when there are two X values corresponding to the entered Y value?**

  In some cases, there are two X values that correspond to a Y value you enter. To understand how Prism handles this situation, you need to know how it interpolates. It starts at the minimum X value and works its way to the largest X value. It stops when it finds the Y value you entered. If it doesn’t find a Y value matching the one you entered, it then looks at X values lower than the minimum shown on the graph and to X values larger than the maximum. If there are two points on the curve that have the Y value you entered as an unknown, Prism will find the lower of the two corresponding X values.

- **Can Prism plot the interpolated values?**

  Sure. The interpolated values are tabulated on a results table. If you want to plot only these values, start from this table, click the New
button, and choose New Graph of Existing data. If you want to add these interpolated values to the graph of the unknowns and its fit, go to that graph, double-click to go to Format Graph, go to the middle (Data on graph) tab, and choose to add the table of interpolated values.

3.2.13 How Prism interpolates

If you ask Prism to compute Y for a given X, it does so by evaluating the equation that defines the curve. That is very straightforward.

If you ask Prism to compute X for a given Y, it is more complicated. Prism does not try to solve the equation algebraically but rather does the interpolation numerically. The results are accurate to at least 6-7 decimal places.

Prism decides on the range of X values to consider. To allow for extrapolation a bit beyond the range of the data, Prism creates an interpolation/extrapolation range that includes the range of the data, and extends in each direction by a distance equal to half the difference between Xmax and Xmin. There are two special cases. When all the data are positive (or zero), that range is clipped to exclude negative numbers. Similarly, when the data are all negative (or zero), the interpolation/extrapolation range is clipped to exclude positive numbers.

Prism then divides that interpolation/extrapolation range into 1000 line segments.

For each value to be interpolated or extrapolated, Prism first tries to interpolate within the range of the X values of the data.

1. It starts with the lowest X value (scans from left to right on the graph). If more than one line segment includes the Y value, Prism only finds the first (lowest X value).

2. Prism then interpolates within that line segment to determine X as accurately as possible. In most cases, it does this by binary bisection. It divides the segment in half and figures out which contains Y. Then it
divides that half in half again. And again. This continues until X is determined as accurately as possible given the numerical precision of the computer. In rare cases, it is possible that Y is not monotonic within the range determined in step 1. In this case, Prism uses linear interpolation rather than binary bisection.

If Prism is not able to interpolate a Y value that corresponds to the entered X value within the X range of the data, it will try to extrapolate a value to the extended range mentioned above. It first looks at X values below the minimum X value in the data, and then at X values above the maximum X value in the data.

When Prism interpolates a value (the X value is in the range of the data), it will also compute a confidence interval if you request it to do so. It does this by determining where the two confidence bands intersect the Y value you entered. When Prism extrapolates a value from the curve, it will not compute a confidence interval for that extrapolated value.

3.2.14 Standard Addition Method

Prism can easily interpolate from a linear or nonlinear standard curve. You perform the assay at a number of known concentrations, fit a line or curve, and interpolate the unknown values.

But there is a problem with interpolating from a standard curve. The results can be incorrect when the unknown sample are contaminated with other substances that alter the assay. This is known as the matrix effect problem.

The Standard Addition Method is a way to bypass this problem. Add various known concentrations (including zero) of known substance to a constant amount of the unknown, and perform the assay. This ensures that all the samples have the same amount of unknown, including any substances that interfere with the assay.

Fit the data with linear regression. The Standard Addition Method only works when the assay measurement (Y) is proportional to the concentration of substance you are measuring.

The dotted line in the figure below shows how the assay results (usually optical density) varies as you change the concentration of the substance you are assaying. The circles and solid line show the assay results in the
presence of an unknown concentration of the substance you are assaying. This pushes the line up and to the left.

The value you want to know is how far the solid line is shifted to the left of the dotted line. There is an easy way to find out: Extrapolate the line down to Y=0. One of the parameters that Prism reports is the X intercept, which will be negative. Take the absolute value, and that is the concentration of the unknown substance. The confidence interval for the X intercept gives you the confidence interval for the concentration of the unknown. Simply multiply both confidence limits by -1.

To plot the data in Prism, you'll want to extend the linear regression line to start at an X value equal to the X intercept (a choice in the Linear regression parameters dialog). You may also want to move the origin to the lower left, a choice on the first tab of the Format Axis dialog. Prism file.

### 3.3 Nonlinear regression tutorials

Prism makes it quite easy to fit a model to your data. If you are new to Prism, choose from the sample XY data sets. These not only show you how to use Prism, but also review the principles of
nonlinear regression, including comparing models, identifying outliers, global fitting, and more.

- **Example: Fitting an enzyme kinetics curve**

- **Example: Comparing two enzyme kinetics models**

- **Example: Automatic outlier elimination (exponential decay)**

- **Example: Global nonlinear regression (dose-response curves)**

- **Example: Ambiguous fit (dose-response)**

- **Example: Interpolating from a sigmoidal standard curve**

### 3.3.1 Example: Fitting an enzyme kinetics curve

1. **Create the data table**

   From the Welcome or New Table dialog, choose to create an XY data table, and select the sample data "Enzyme kinetics -- Michaelis-Menten".

2. **Inspect the data**

   The sample data will be partly covered by a floating note explaining how to fit the data (for people who are not reading this help page). You can move the floating note out of the way, or minimize it.
The data are in triplicate. Some values are missing, and Prism always handles these just fine.

3. View the graph

Prism automatically created a graph and gave it the same name as the data table. Click on the Michaelis-Menten graph in the graphs section.

Since this is the first time you are viewing the graph, Prism will pop up the Change Graph Type dialog. Select the fifth choice, to plot individual replicates rather than mean and error bars.

The graph Prism makes automatically is fairly complete. You can customize the symbols, colors, axis labels, position of legend, etc.

4. Choose nonlinear regression

Click the Analyze button and choose Nonlinear regression from the list of XY analyses.

Even faster, click the shortcut button for nonlinear regression.
5. Choose a model

On the Fit tab of the nonlinear regression dialog, open the equation folder, Enzyme Kinetics - Substrate vs. Velocity. Then choose the Michaelis-Menten equation.

Learn more about the principles of enzyme kinetics and about fitting Michaelis-Menten curves.

For this example, leave all the other settings to their default values.

Click OK to see the curves superimposed on the graph.
6. Inspect the graph

![Michaelis-Menten data graph]

7. Inspect the results

<table>
<thead>
<tr>
<th>Nonlin fit</th>
<th>A</th>
<th>Enzyme Activity</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td></td>
<td>Y</td>
</tr>
<tr>
<td>2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>VMAX</td>
<td>1353</td>
</tr>
<tr>
<td>4</td>
<td>KM</td>
<td>5.886</td>
</tr>
<tr>
<td>5</td>
<td>Std. Error</td>
<td>75.93</td>
</tr>
<tr>
<td>6</td>
<td>VMAX</td>
<td>0.8498</td>
</tr>
<tr>
<td>7</td>
<td>KM</td>
<td>1197 to 1509</td>
</tr>
<tr>
<td>8</td>
<td>95% Confidence Intervals</td>
<td>3.933 to 7.839</td>
</tr>
</tbody>
</table>

The goal of nonlinear regression is to find the best-fit values of the parameters. These are reported at the top of the table. You can't really interpret the best-fit values without knowing how precise they are, and this is reported both as standard errors and confidence intervals.
8. Go back and perform the replicates test

The replicates test assesses the adequacy of the fit by comparing the scatter among the triplicates with the scatter of points around the curve. It is not calculated by default, so the results do not appear in the results of step 7.

You don't have to do the fit over again. Instead click the button in the upper left corner of the results table to return to the nonlinear regression dialog.

Go to the Diagnostics tab, and check the option to perform the replicates test. Note that you can also check an option to make your settings here become the default for future fits.
The P value is small (0.013). This means that the scatter of the data from the curve is greater than you’d expect from the variation among triplicates. This suggests that you might want to consider fitting an alternative model, which we do in the next example.
3.3.2 Example: Comparing two enzyme kinetics models

1. Open analysis parameters dialog

This example continues from the previous one. Click the analysis parameter button again.

2. Choose the first model

On the Fit tab, choose (or make sure you still have chosen) the Michaelis-Menten equation.
3. Choose the other model

Go to the Compare tab.

Choose: For each data set, which of two equations (models) fits best?

There are two ways to compare models. For this example, choose the extra sum-of-squares F test.

For the second equation, choose "Allosteric sigmoidal" from the Enzyme kinetics section.
4. View the results of the model comparison

The top part of the results summarizes the comparison. The P value is low, suggesting that the simpler (Michaelis-Menten) model is too simple and should be rejected. The allosteric model fits significantly better.

5. View the numerical results of the fit to the allosteric model

Scroll down to the best-fit parameter values for the allosteric model.
The P value of the replicates test is high, which means the scatter of points around the curve is consistent with variability of replicates from each other.

The parameter H equals 2.0, with a 95% confidence interval ranging from 1.5 to 2.5. A value of 2.0 suggests that this enzyme might be a dimer. When H equals 1.0, the allosteric model is identical to the Michaelis-Menten model.

Further interpretation must, of course, be in the context of what is known about this enzyme from prior work. Statistics is only part of analyzing scientific data.

6. View the graph

Since the allosteric model fit significantly better, that is the model that Prism uses when it plots a curve on the graph. You can barely see that it has a sigmoidal shape. You can also see that you should collect more data with substrate concentrations between 0 and 5 to fully define this curve.
3.3.3 Example: Automatic outlier elimination (exponential decay)

Prism implements a unique method to identify outliers when fitting curves with nonlinear regression. This example shows you how easy it is to identify outliers with Prism. Learn more about how this method works, when it is useful, and when it should be avoided.

1. Create the data table

From the Welcome or New Table dialog, choose to create XY data table, and select the sample data "Eliminating outliers during nonlinear regression".

2. Inspect the data table and graph

The data table has values for two data sets (control and treated) with triplicate subcolumns.

Since this is the first time you are viewing the graph, Prism will pop up the Change Graph Type dialog. Select the fifth choice, to plot individual replicates rather than mean and error bars.
3. Choose nonlinear regression

Click the Analyze button and choose Nonlinear regression from the list of XY analyses.

Alternatively, click the shortcut button for nonlinear regression.

4. Choose a model, and choose automatic outlier detection

On the Fit tab of the nonlinear regression dialog, open the panel of exponential equations and choose: One phase decay.

Also choose automatic outlier elimination. Learn about when outlier elimination is helpful, when it should be avoided, and how it works.
5. View the graph

Prism identified the outlier, and plotted it in red, overlaid on top of the data graph. After identifying the outlier, Prism fit the remaining data points as if the outlier wasn't present. Before accepting the results, think about why the point was an outlier. Remember, not all outliers are "bad" points.

Double click on the graph to bring up the Format Graph dialog. Go to the second tab. You can see that this graph now has three data sets, the data, the curve fit, and the outliers. Read more about graphing outliers.

3.3.4 Example: Global nonlinear regression (dose-response curves)

1. Create the data table

From the Welcome or New Table dialog, choose to create an XY data table, and select the sample data "Dose-response: EC50 shift by global fitting". Note that this choice is in a drop down menu of sample data sets.
2. Inspect the data

The sample data may be partly covered by a floating note explaining how to fit the data (for people who are not reading this help page). You can move the floating note out of the way, or minimize it.

<table>
<thead>
<tr>
<th>X</th>
<th>A:Y1</th>
<th>A:Y2</th>
<th>B:Y1</th>
<th>B:Y2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>-7.5</td>
<td>341</td>
<td>298</td>
<td>295</td>
</tr>
<tr>
<td>2</td>
<td>-7.0</td>
<td>671</td>
<td>752</td>
<td>616</td>
</tr>
<tr>
<td>3</td>
<td>-6.5</td>
<td>874</td>
<td>721</td>
<td>362</td>
</tr>
<tr>
<td>4</td>
<td>-6.0</td>
<td>1000</td>
<td>951</td>
<td>444</td>
</tr>
<tr>
<td>5</td>
<td>-5.5</td>
<td>1305</td>
<td>1265</td>
<td>882</td>
</tr>
<tr>
<td>6</td>
<td>-5.0</td>
<td>1254</td>
<td>1351</td>
<td>1354</td>
</tr>
<tr>
<td>7</td>
<td>-4.5</td>
<td>1265</td>
<td>1411</td>
<td>1452</td>
</tr>
</tbody>
</table>

The X values are the logarithm of the concentration of agonist. The Y values are responses, in duplicate, in two conditions.

3. View the graph

Since this is the first time you are viewing the graph, Prism will pop up the Change Graph Type dialog. Select the first choice, to plot mean and error bars, and choose the kind of error bars you prefer.

4. Choose nonlinear regression

Click  and choose Nonlinear regression from the list of XY analyses.
Alternatively, click the shortcut button for nonlinear regression.

5. Choose a model

On the Fit tab of the nonlinear regression dialog, open the panel of inhibitory dose-response models and choose: log(inhibitor) vs. response -- variable slope.

For now, leave all the other settings to their default values.

Click OK to see the curves superimposed on the graph.
6. Inspect the results

The control results are labeled ambiguous. This means that Prism is unable to find a unique curve through the data. Lots of other sets of parameter values would lead to curves that fit just as well. You can see which parameters are ambiguous by looking at the 95% confidence intervals. Instead of reporting an interval, Prism reports 'very wide' for the Bottom and logEC50.

The data do not define a bottom plateau for the control (circles) data set, so its best-fit value is ambiguous. The EC50 is the concentration that gives a response half way between the bottom and top plateaus of the curve. If the bottom is ambiguous, so is the EC50.

The treated curve is not labeled 'ambiguous', but the confidence intervals are wider than you’d like.
7. Go back to the dialog, and share three parameters

You can get much better results from this data set if you are willing to assume that that the top and bottom plateaus, and the slope, are the same under control and treated conditions. In other words, you assume that the treatment shifts the EC50 but doesn't change the basal response, the maximum response, or the Hill slope.

Return to the nonlinear regression dialog by clicking the button in the upper left of the results table.

Go to the constraints tab and choose to share the value of Bottom, Top, and HillSlope. When you share these parameters, Prism fits the data sets globally to find one best-fit value for Bottom, Top and HillSlope (for both data sets) and separate best-fit values for the logEC50.

8. View the revised graph and results
The fit is no longer labeled ‘ambiguous’ and the confidence intervals are much tighter.

9. Statistically compare the two logEC50 values

Go back to the parameters dialog for nonlinear regression and go to the Compare tab. Check the option to test whether one curve adequately fits all the data sets.
Prism will now fit the data two ways. The first is the same as before, fitting a separate IC50 for each data set. The second fit shares all the parameters. In this case, three parameters were already shared but one wasn’t. So in this second fit, all four parameters are shared, so Prism fits one curve through all the data, ignoring which treatment group they are in. It compares the sum-of-squares (really the sum-of-sum-of squares since there are two data sets fit in each case), using the extra sum-of-squares F test. The results are shown at the top of the results sheet.
The P value is tiny, so we reject the null hypothesis that the two IC50 values are identical in the population, and instead conclude that the two IC50 values are different.

10. Fit the ratio of two IC50 values directly

From the results in step 8, you can compute what you want to know -- the ratio of the two EC50 values.

But Prism can calculate this value directly.

Go back to the analysis parameters dialog, and on the Fit tab, change the equation to "EC50 shift" from the "Dose-response - Special" group of equations. Accept all defaults and click OK. The graph will look identical, as the model is equivalent. But now, rather than fitting two logEC50 values, Prism fits one and also fits the ratio.

This equation was designed to do exactly what is needed for this example. Read about how this equation was set up, so you can construct your own equations when necessary.
3.3.5 Example: Ambiguous fit (dose-response)

1. Create the data table

From the Welcome or New Table dialog, choose to create XY data table, select the sample data "Dose-Response - Ambiguous until constrained". Note that this choice is in a drop down menu of sample data sets.

2. Inspect the data table

The X values are the logarithms of molar concentration. The Y values are responses, entered as mean and SD. With Prism, you can either enter replicate values or enter error values (here SD) computed elsewhere.

<table>
<thead>
<tr>
<th>Title</th>
<th>X</th>
<th>Mean</th>
<th>SD</th>
<th>N</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>log(agonist)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>-7.0</td>
<td>135</td>
<td>24</td>
<td>3</td>
</tr>
<tr>
<td>2</td>
<td>-6.5</td>
<td>258</td>
<td>20</td>
<td>3</td>
</tr>
<tr>
<td>3</td>
<td>-6.0</td>
<td>322</td>
<td>19</td>
<td>2</td>
</tr>
<tr>
<td>4</td>
<td>-5.5</td>
<td>398</td>
<td>35</td>
<td>4</td>
</tr>
<tr>
<td>5</td>
<td>-5.0</td>
<td>458</td>
<td>21</td>
<td>3</td>
</tr>
<tr>
<td>6</td>
<td>-4.5</td>
<td>509</td>
<td>26</td>
<td>3</td>
</tr>
<tr>
<td>7</td>
<td>-4.0</td>
<td>512</td>
<td>31</td>
<td>3</td>
</tr>
</tbody>
</table>

3. View the graph

Since this is the first time you are viewing the graph, Prism will pop up the Change Graph Type dialog. Select the first choice, to plot mean and error bars, and choose the kind of error bars you prefer.
4. **Choose nonlinear regression**

Click **Analyze** and choose from the list of XY analyses.

Even faster, click the shortcut button for nonlinear regression.

5. **Choose a model**

On the Fit tab of the nonlinear regression dialog, open the panel of Stimulatory dose-response equations and choose: log(agonist) vs. response -- Variable slope.
Learn more about fitting dose-response curves. For this example, leave all the other settings to their default values. Click OK to see the curves superimposed on the graph.
6. Inspect the graph

The curve goes through the point nicely, and looks fine.

7. Inspect the results

Go to the results page, and view the results.
Note the word 'ambiguous' at the top of the results. This means that Prism was unable to find a unique fit to these data. Lots of sets of parameter values would lead to curves that fit just as well. Learn more about ambiguous fits.

Prism does not report confidence intervals for the logEC50 or the Bottom of the curve, but instead simply says the intervals are 'very wide'. That tells you it was impossible to fit those parameters precisely.

While the Y values of the data range about 100 to about 600, the best-fit value for the bottom of the curve is -1067. Furthermore, the best-fit value of the logEC50 is outside the range of the data.

Even though the curve is close to the points (the $R^2$ is 0.9669), the best-fit parameter values are useless.
8. Constrain the Bottom of the curve

The problem is simple. You have asked Prism to find best-fit value for four parameters representing the bottom and top plateaus of the curve as well as the mid point and steepness. But the data simply don't define the bottom of the curve. In fact, Prism the best fit value of Bottom is very very far from the data.

These data were calculated so the basal nonspecific response was subtracted away. This means that you know that the response (Y) at very low concentrations of agonist (very low values of X) has to be zero. Prism needs to know this to fit the data sensibly.

You don't have to do the fit over again. Instead click the button in the upper left corner of the results table to return to the nonlinear regression dialog.

Go to the Constrain tab, and check constrain the parameter Bottom to have a constant value which you set to 0.0.

9. Inspect the revised results

Now the results make sense. The logEC50 is in the middle of the range of X values. The confidence intervals are reasonably tight. And, of course, the results are no longer 'ambiguous'.
3.4 Nonlinear regression with Prism

Prism's nonlinear regression dialog has seven tabs, explained here. It can seem overwhelming at first. But you don't have to learn about all the choices when you first fit a curve. Start by focusing on the choices in the first (Fit) tab, and then learn about the other choices later.

- Before learning all the details, first understand the big picture -- the basic steps required to fit a model to data.

- So many choices! Which ones are essential?
3.4.1 How to fit a model with Prism

**Step 1. Enter data**

Create an XY table, and enter data. If you have replicate Y values at each X value, format the table for entry of replicates.

From an XY table or graph, click the shortcut button to fit a model with nonlinear regression. Or click Analyze and select from the analyze dialog.

**Step 2. Choose a model**

Nonlinear regression fits a model to your data. You must, therefore, choose a model or enter a new model.

Why a computer program cannot pick a model for you.

**Step 3. Choose (or review) initial values**

Nonlinear regression is an iterative procedure. The program must start with estimated values for each variable that are in the right "ball park" - say within a factor of five of the actual value. It then adjusts these initial values to improve the fit. See How nonlinear regression works.

GraphPad Prism provides initial values automatically if you use a built-in equation. If you enter your own equation, you can also enter rules for initial values. For example the initial value of one parameter may be twice the maximum Y value in the data, while the initial value of another parameter may equal the average of the highest and lowest X values. Once you define these rules, Prism will compute appropriate initial values based on the range of your data.

The Initial Values tab of the Nonlinear regression dialog lets you review, and override, the computed initial values.
You'll find it easy to estimate initial values if you have looked at a graph of the data, understand the model, and understand the meaning of all the parameters in the equation. Remember that you just need an estimate. It doesn't have to be very accurate. If you are having problems estimating initial values, set aside your data and simulate curves using the model. Change the variables one at a time, and see how they influence the shape of the curve. Once you have a better feel for how the parameters influence the curve, you might find it easier to estimate initial values.

When fitting a simple model to clean data, it won't matter much if the initial values are fairly far from the correct values. You'll get the same best-fit curve no matter what initial values you use, unless the initial values are extremely far from correct. Initial values matter more when your data have a lot of scatter or your model has many variables.

**Step 4. Decide whether to constrain any parameters**

When performing nonlinear regression, you don't have to fit each parameter in the equation. Instead, you may fix one or more of the parameters to constant values. It is often helpful to define constants when you have only a few data points. For example, you might fix the bottom plateau of a sigmoid curve or exponential decay to zero.

Remember that nonlinear regression programs have no "common sense". You need to think about how you did the experiment, and decide whether some of the parameters should be fixed. For example, if a background signal has already been subtracted, it makes sense to fix the bottom plateau of a dose-response curve or an exponential decay curve to zero.

Prism also let you constrain parameters to a certain range of values.

**Step 5. If you are fitting two or more data sets at once, decide whether to share any parameters**

If you enter data into two or more data set columns, Prism will fit them all in one analysis. But each fit will be independent of the others unless you specify that one or more parameters are shared. When you share parameters, the analysis is called a **global nonlinear regression**.
Step 6. Decide on a weighting scheme

Nonlinear regression programs generally weight each point equally. But there are many ways to differentially weight the points.

Step 7. Choose other options

Read about choices on the Range, Output and Diagnostics tab.

3.4.2 Which choices are essential?

So many choices! Prism's nonlinear regression dialog has seven tabs, and can seem overwhelming at first. But you don't have to learn about all the choices when you first fit a curve.

If your goal is to fit a standard curve from which to interpolate unknown values

If your goal is to interpolate from a standard curve, you can ignore most of the choices that Prism offers. You do need to pick a model, but you should judge the adequacy of the model visually. If it goes through the points without wiggling too much, it is fine for interpolation. Note that Prism 6 offers a simplified curve fitting analysis for interpolation.

If your goal is to fit a model to determine best-fit parameters

If your goal is to fit a model to determine parameter values, then choosing a model is the essential first step. You should not expect a computer (or a software company's tech support) to choose a model for you. Choosing a model, and deciding which parameters should be constrained and which should be shared among data sets, is a scientific decision that is fundamental to analyzing your data. The other choices are useful, but can be put aside when you are first learning curve fitting.

3.4.3 Nonlinear regression choices

- Choose the model (equation) on the Fit tab. This is also where you choose to eliminate outliers and to interpolate values from the fit curve.
• **Compare tab**. Choose to compare the fit of two models to each data set, or to compare the fit of two (or more) data sets.

• **Constrain tab**. Fix parameter values to constant values and/or designate parameters to be shared among all data sets.

• There are many situations where you don't want to weight all points equally. Set unequal weighting on the **Weights tab**.

• Nonlinear regression is an iterative procedure. The program must start with estimated initial values for each parameter. It then adjusts these initial values to improve the fit. View those values on the **Initial values tab**.

• **Range tab**. Set the X range of data to fit, the starting and ending X values for the plotted curve, and whether you want to view a table of XY coordinates that define the curve.

• **Output tab**

• **Diagnostics tab**. Prism can create more detailed results tables.

### 3.4.3.1 Fit tab

**Choose an equation**

Selecting an equation is the most important step in fitting a curve. The choice **cannot be automated**.

If your goal is to fit a model in order to understand your data and make comparisons, then choosing a model is a scientific decision that you must make with care. If your goal is to interpolate unknowns from a standard curve, then it matters less which equation you pick, so long as it ends up creating a smooth curve through your data.

Some tips:

• Prism provides a long list of equations that you can choose. But if these don't fit your needs, don't be afraid to **create or clone an equation** to fit your needs.
• Part of choosing a model is choosing constraints. Don't skip that step. For example, if you choose a sigmoidal dose response model, you must decide whether you wish Prism to find the best fit value of the bottom plateau based on the trend of the data. The alternative is to constrain the bottom plateau to equal zero, if you have subtracted off a baseline, or some other value (defined by controls). A computer can't make these decisions for you. Choosing which constraints to apply to your model is a fundamental decision in data analysis that can have a huge impact on the results.

• If you are fitting several data sets at once, part of choosing a model is deciding which parameters you want to share between data sets. When you share a parameter (a choice on the Constrain tab), Prism finds one best-fit value for the parameter that applies to all the data sets. Read more about shared parameters (global fitting).

Fitting method

Prism offers three methods to fit curves.

- Least squares (ordinary fit)
- Robust fit
- Automatic outlier elimination

If you aren't sure which method to choose, pick least-squares regression, as it is standard. Robust regression is less affected by outliers, but it cannot generate confidence intervals for the parameters, so has limited usefulness. Automatic outlier removal is extremely useful, but can lead to invalid (and misleading) results in some situations, so should be used with caution.

Interpolate

Check this option in order to interpolate the concentration of unknown samples from the best-fit curve. Learn more.

With this option, Prism will report the Y value for any X values you enter, and the X value for any Y values you enter (including extrapolating in each direction a distance equal to half the length of the X axis).
3.4.3.2 Compare tab

What question are you asking?

When fitting biological data with regression, your main objective is often to discriminate between different models, to ask if an experimental intervention changed a parameter, or to ask if the best-fit value of a parameter differs significantly from a theoretical value. Learn more about these four kinds of comparisons. Your choice, of course, has to be based on your experimental goals.

Comparison method

Prism can perform the comparison using two alternative methods: the extra sum-of-squares F test, and using Akaike’s information criteria. Use these guidelines to choose:

- In most cases, the two models will be 'nested'. This means that one model is a simpler case of the other. For example, a one-phase exponential model is a simpler case of a two-phase exponential model. Either the F test or the AICc method may be used with nested models. The choice is usually a matter of personal preference and tradition. Basic scientists in pharmacology and physiology tend to use the F test. Scientists in fields like ecology and population biology tend to use AICc.
• If the models are not nested, then the F test is not valid so you should choose AICc. Note that Prism does not enforce this. It will calculate the F test even if the models are not nested, but the results won't be useful.

The extra sum-of-squares F test only makes sense when the models being compared have different numbers of parameters, and so have different numbers of degrees of freedom. If you want to compare two models with the same number of parameters, choose the AIC method.

Check the option on the right, and before running the extra sum-of-squares F or AICc comparison, Prism checks whether both models actually fit the data. If one fit is ambiguous or flagged, Prism will choose the other without any calculation.

If you are comparing the fits of three or more models, fit individually, you can ask Prism to report the AICc with each fit. This is a choice on the Diagnostics tab.

3.4.3.3 Constrain tab

The importance of constraints

Prism lets you constrain each parameter to a constant value, constrain to a range of values, share among data sets (global fit), or define a parameter to be a column constant. This is an important decision, and will influence the results.

Constrain to a constant value

You won't always want to fit all the parameters in a model. Instead, you can fix one or more parameters to constant values. For example, if you have normalized a dose-response curve to run from 0 to 100, constrain Top to equal 100 and Bottom to 0.0. Similarly, if you have subtracted a
baseline so you know that the exponential decay curve has to plateau at Y=0.0, you can constrain the Bottom parameter to equal 0.0.

Deciding to constrain a parameter to a constant value can make a huge difference in the results. Remember Prism has no common sense. It doesn't know if a curve has to plateau at zero unless you tell it.

**Constrain to a range of values**

Constrain to a range of values to prevent Prism from letting parameters take on impossible values. For example, you should constrain rate constants to only have values greater than 0.0, and fractions (say the fraction of binding sites that are high affinity) that have a value between 0.0 and 1.0. Setting this kind of constraint can have four outcomes:

- The constraint is irrelevant, as the parameter never would have taken on a value in the forbidden range.

- The constraint helped speed up the fit. Nonlinear regression works by iteratively changing the values of the parameters. With some complicated fits, the nonlinear regression process can 'get confused' and end up spending time exploring parameter values that make no sense. Constraining the values of one or more parameters can prevent the nonlinear regression process from being led astray. With huge numbers of data points, you might see a noticeable speeding up of the fitting process.

- The constraint helped nonlinear regression choose from several local minima. Nonlinear regression works by changing parameter values step by step until no small change affects the sum-of-squares (which quantifies goodness-of-fit). With some models, there can be two sets of parameter values that lead to local minima in sum-of-squares. Applying a constraint can ensure that nonlinear regression finds the minimum with scientifically relevant values, and ignores another minimum that fits the curve well but using parameter values that make no scientific sense (i.e. negative concentrations).

- The constraint prevents nonlinear regression from finding a minimum sum-of-squares. Instead, the best the program can do (while obeying the constraint) is set the parameter to the limit of the constrained range. Prism then reports that the fit 'hit constraint'.

In the first case, the constraint is harmless but useless.
In the next two cases, the constraint helps the nonlinear regression reach sensible results. Essentially, the constraint can give the nonlinear regression process some scientific judgment about which parameter values are simply impossible. These cases are really what constraints are for.

The last case, when the fit ends with a parameter set to one end of its constraint, it is tricky to interpret the results. [46]

Sharing parameters among data sets. Global nonlinear regression.

If you are fitting a family of curves, rather than just one, you can choose to share some parameters between data sets. For each shared parameter, Prism finds one (global) best-fit value that applies to all the data sets. For each non-shared parameter, the program finds a separate (local) best-fit value for each data set. Global fitting is a very useful tool in two situations:

- The parameter(s) you care about are determined from the relationship between several data sets. Learn more. [48]
- Each dataset is incomplete, but the entire family of datasets defines the parameters. See example. [47]

Data set constant

When you fit a family of curves at once, you can set one of the parameters to be a data set constant. Prism offers two data set constants.

**Column title.** The value comes from the column title, which can be different for every data set. This parameter becomes almost a second independent variable. It has a constant value within any one data set, but a different value for each data set. For example, when fitting a family of enzyme progress curves in the presence of various concentrations of inhibitor, the inhibitor concentration can be entered into the column title of the data table. View an example [46].

**Mean X.** The value is the mean of all the X values for which there are Y values in that data set. This is used in centered polynomial regression.
Different constants for different data sets

When you use the Constrain tab to set a parameter to a constant value, you do so for all data sets. It is not possible to enter a different constant value for each data set (unless you use the column title as the constant value as described in the previous section. If you want a parameter to have a different constant value (not the column title) for each data set, you'll need to write a user defined equation, and use special notation to assign a different value for each data set. You'll need to hard code the constants within the user-defined equation, and cannot choose them in the nonlinear regression parameters dialog. For example:

\[
\text{Y} = \text{Bottom} + \text{span} \times (1 - \exp(-1 \times \text{K} \times \text{X}))
\]

In this example, the parameter Bottom is set to 4.5 when fitting data set A, to 34.5 when fitting data set B, and to 45.6 when fitting data set C.

3.4.3.4 Weights tab

 ROUT coefficient

If you ask Prism to automatically exclude outliers (a choice on the Fit tab) or to count (but not remove from the analysis) outliers (a choice on the Diagnostics tab), set the ROUT coefficient to determine how aggressively Prism defines outliers.

We recommend using a value of 1%. Our simulations have shown that if all the scatter is Gaussian, Prism will falsely find one or more outliers in about 2-3% of experiments. If there really are outliers present in the data, Prism will detect them with a False Discovery Rate less than 1%. See reference 1.

If you set Q to a higher value, the threshold for defining outliers is less strict. This means that Prism will have more power to detect outliers, but also will falsely detect 'outliers' more often. If you set Q to a lower value, the threshold for defining outliers is stricter. This means that Prism will have a less power to detect real outliers, but also have a smaller chance of falsely defining a point to be an outlier.
If you set Q to 0, Prism will fit the data using ordinary nonlinear regression without outlier identification.

**Unequal weighting**

It is often useful to differentially weight the data points. [Learn why](#).

Prism offers seven choices on the Weight tab of nonlinear regression:

- **No weighting.** Regression is most often done by minimizing the sum-of-squares of the vertical distances of the data from the line or curve. Points further from the curve contribute more to the sum-of-squares. Points close to the curve contribute little. This makes sense, when you expect experimental scatter to be the same, on average, in all parts of the curve.

- **Weight by 1/Y^2.** In many experimental situations, you expect the average distance (or rather the average absolute value of the distance) of the points from the curve to be higher when Y is higher. The points with the larger scatter will have much larger sum-of-squares and thus dominate the calculations. If you expect the relative distance (residual divided by the height of the curve) to be consistent, then you should weight by 1/Y^2.

- **Weight by 1/Y.** This choice is useful when the scatter follows a Poisson distribution -- when Y represents the number of objects in a defined space or the number of events in a defined interval.

- **Weight by 1/Y^k.** Also called "General weighting". [Read more](#).

- **Weight by 1/X or 1/X^2.** These choices are used rarely. Only choose these weighting schemes when it is the standard in your field, such as a linear fit of a bioassay.

- **Weight by 1/SD^2.** If you enter replicate Y values at each X (say triplicates), it is tempting to weight points by the scatter of the replicates, giving a point less weight when the triplicates are far apart so the standard deviation (SD) is high. But unless you have lots of replicates, this doesn't help much. The triplicates constituting one mean could be far apart by chance, yet that mean may be as accurate as the others. Weighting needs to be based on systematic changes in scatter. The choice to weight by 1/SD^2 is most useful when you want to use a weighting scheme not available in Prism. In this case, enter data as
mean and SD, but enter as "SD" weighting values that you computed elsewhere for that point. In other words, the values you enter in the SD subcolumn are not actually standard deviations, but are weighting factors computed elsewhere.

**Notes on weighting**

- If you have normalized your data, **weighting rarely makes sense**.
- **Simulations can show** you how much difference it makes if you choose the wrong weighting scheme.
- If you choose unequal weighting, **Prism takes this into account** when plotting residuals.
- Prism accounts for weighting **when it computes R** when it computes R.
- Learn about the **math of weighting** and how **Prism does the weighting**.
- If you choose robust regression (on the first, Fit, tab) then choices you make on the weighting tab will not influence the fitting, so will no influence the best-fit parameter values or the curve. However your choice of weighting will have an impact on the residuals Prism computes and graphs and on how it identifies outliers. Both these choices (residuals, and outlier counting) are on the Diagnostics tab of nonlinear regression.

**Replicates**

- Consider each replicate Y value as an individual point.
- Only consider the mean Y value of each point.

In most experiments, it is fair to consider each replicate to be an independent data point, and you should choose that first choice unless you have a strong reason not to.

If you entered the data as mean, n, and SD or SEM the wording is a bit different. You can account for sd and n and get exactly the same results you would have gotten had you entered raw data (except for robust
regression, and outlier determination which require raw data). Or you can fit the means only, and ignore n and the entered SD or SEM.

Here is an example where you would want to fit only the means: You performed a dose-response experiment, using a different animal at each dose with triplicate measurements. The three measurements are not independent. If one animal happens to respond more than the others, that will affect all the replicates. Since the replicates are not independent, you should fit the means.

Reference


3.4.3.5 Initial values tab

Why initial values?

Nonlinear regression is an iterative procedure. The program must start with estimated initial values for each parameter. It then adjusts these initial values to improve the fit.

How does Prism provide initial values automatically?

Every equation built-in to Prism, as well as equations you define, include rules to compute initial values. These rules use the range of the X and Y values to come up with initial values, which become the original automatic initial values. You can change the rules for user-defined equations, and can clone built-in equations to make them user-defined. The new rules will be invoked when you next choose this equation for a new analysis. It won't change the initial values for the analysis you are working on.

Are the initial values reasonable?

If nonlinear regression doesn't report any results, or if they seem wrong, the problem might be bad initial values. To check whether the initial values are reasonable, check "Don't fit the curve. Instead plot the curve defined by initial values." on the Diagnostics tab. When you click OK from the nonlinear regression dialog, Prism will not fit a curve but will instead generate a curve based on your initial values. If this curve
is not generally in the vicinity of the data points, change the initial values before running nonlinear regression.

**Changing the initial values**

In most cases, the automatic initial values will work fine. If you have trouble getting a sensible fit, you might need to tweak the initial values. You can override the automatic initial values for each parameter for each data set:

1. Select a data set on the top part of the initial values tab. Or hold down the Ctrl key and select several data sets. Or click "Select all" in the top right of the tab.

2. Uncheck "choose automatically" and enter the new initial value.

You will find it easy to estimate initial values if you have looked at a graph of the data, understand the model, and understand the meaning of all the parameters in the equation. Remember that you just need an estimate. It doesn't have to be very accurate.

**How much difference do initial values make?**

When fitting a polynomial model, it makes no difference what values you enter as initial values. When fitting other models, the importance of the initial values depends on how well the data define the curve and on how many parameters are in the model. Initial values matter the most when your data have a lot of scatter and don't define the model very well, and your model has many parameters.

### 3.4.3.6 Range tab

**Ignore points outside of a specified X range**

If you choose these options nonlinear regression will completely ignore all data where the X value is less than, or larger than, the thresholds you set. This could be useful if you collect data over time, and only want to fit data within a certain range of time points.

**Define the curve**

In addition to fitting the model to your data, Prism also superimposes the curve on your graph. Choose where the curves should begin and end.
Table of XY coordinates

Check this option if you want to see a table of XY coordinates of line segments defining the curve.

In prior versions of Prism, this table was critical to plotting the curve and interpolating unknowns. But Prism 5 can graph the curve and interpolate unknowns from the curve, without creating this table. Check the option to create this table only if you want to copy (or export) the table into another program. In most cases, you will have no need to view this table, and so should leave the option unchecked.

3.4.3.7 Output tab

Summary table

When analyzing several data sets, the results table is rather lengthy. To display key results on a summary table, check the option box to create a summary table and select the variable you wish to summarize. Prism creates a summary table (as an additional results view) that shows the best-fit value of that parameter for each data set, and graphs this table.

Depending on your choices in the dialog, this may be a bar graph or an XY graph. It shows the best-fit value of a selected parameter for each data set on the table. In some cases, you may analyze the summary table with linear or nonlinear regression. For example, the summary graph may show the best-fit value of a rate constant as a function of concentration (obtained from the column titles of the original data). You can fit a line or curve to that graph.

When Prism compares the fits of two equations, it shows only the results for the second equation. Since this may not be helpful, we suggest that you only make summary tables when fitting a single equation.

Number of digits in output

Choose how many significant digits you want to see in the main results table. This is especially useful if you embed the results table on a graph or layout. If you choose to show many digits of precision in the results, you probably should also choose strict convergence criteria on the Diagnostics tab.
For the table of residuals and the curve itself, you can choose a number format by selecting column(s) and bring up the Decimal Format dialog.

**Additional output (legacy for old Prism versions)**

The options here (dose ratios, and Ki from IC50) are available only to be compatible with earlier versions of Prism (so Prism can open files made with older versions). They are almost always unavailable, becoming available only if you select certain equations from the classic equations list.

Since Prism 5, you can calculate a Gaddum/Schild EC50 shift directly, without need to separately compute dose ratios. Similarly, you can fit competitive binding curves directly to determine the Ki for one or two sites, without a separate Cheng-Prusoff calculation of Ki from IC50(1).


**3.4.3.8 Confidence tab**

**Confidence intervals of the parameters**

If you reason to run nonlinear regression is to interpolate unknown values, you won't really care about the values of the parameters so won't care about the confidence interval of the parameters. But if you do care about the values of the parameters, we suggest that you always ask Prism to report confidence intervals, as inspecting the confidence intervals of best-fit parameters is an essential part of evaluating any nonlinear fit.

**How to compute the intervals**

Prism offers two methods to compute the confidence intervals.
• Asymptotic approximate symmetrical confidence intervals. These were the only confidence intervals reported by Prism 6 and earlier and by most programs. But since the true uncertainty in the parameter's value is often asymmetrical, these symmetrical intervals are not always accurate. We suggest choosing them only when you need to compare Prism's results with other programs, when you need to be consistent with earlier work, or when you have so much data that the profile likelihood method is too slow.

• Asymmetrical (and thus more accurate) profile likelihood confidence intervals (new to Prism 7). These do a better job of quantifying how precisely you know the parameters value, so we suggest using them routinely. The only disadvantage is that the calculations are more complex so might be noticeably slower with huge data sets (especially with user-defined equations).

**How to report the intervals**

Prism can report the confidence intervals in two ways: as a range or as separate blocks of lower and upper confidence limits (useful if you want to paste the results into another program). The former is easier to read. The latter might be better if you are tabulating the results elsewhere.
Confidence and prediction bands

The 95% confidence bands enclose the area that you can be 95% sure contains the true curve. It gives you a visual sense of how well your data define the best-fit curve.

The 95% prediction bands enclose the area that you expect to enclose 95% of future data points. This includes both the uncertainty in the true position of the curve (enclosed by the confidence bands), and also accounts for scatter of data around the curve. Therefore, prediction bands are always wider than confidence bands. When you have lots of data points, the discrepancy is huge.

Learn more about confidence and prediction bands.

Standard errors of the parameters

Standard errors are intermediate values used to compute the symmetrical confidence intervals, but are not very useful by themselves.

You may want to include standard errors in the results to compare Prism's results to those of another program that doesn't report confidence intervals or to collaborate with colleagues who don't understand confidence intervals. But we recommend routinely turning off reporting of the standard errors, because they do a poor job of conveying the precision of the best-fit parameter values.
3.4.3.9 Diagnostics tab

**Do the initial parameter values define a curve near the data?**

- **Don't fit the curve.** Instead plot the curve defined by the initial values of the parameters.
- **Fit the curve.**

Nonlinear regression works iteratively, and begins with initial values for each parameter. Check "don't fit the curve" to see the curve generated by your initial values. If the curve is far from the data, go back to the initial parameters tab and enter better values for the initial values. Repeat until the curve is near the points. Then go back to the Diagnostics tab and check "Fit the curve". This is often the best way to diagnose problems with nonlinear regression.

While fitting a curve, Prism will stop after the maximum number of iterations set here. If you are running a script to automatically analyze many data tables, you might want to lower the maximum number of iterations so Prism won't waste time trying to fit impossible data.

**How to quantify the standard deviation of the residuals?**

The standard way to report the standard deviation of the residuals is with the Sy.x also called Se. Prism can also report the root mean square error, but we don't recommend it. How do the values differ?

**How to quantify goodness-of-fit.**

- **R**
- **Adjusted R**
- **Sum-of-Squares**
- **Sy.x**
- **RMSE**
- **AICc**

**R** and **Adjusted R**

You will probably want to ask Prism to report **R**, simply because it is standard to do so, even though knowing **R** doesn't really help you interpret the results. Reporting the sum-of-squares and **Sy.x** will only be useful if you want to compare Prism's results to those of another program, or you want to do additional calculations by hand.

The **adjusted R** takes into account the number of parameters fit to the data, so has a lower value than **R** (unless you fit only one
Regression with Prism

How to quantify the standard deviation of the residuals?

The standard way to report the standard deviation of the residuals is with the $\text{Sy.x}$ also called $\text{Se}$. Prism can also report the root mean square error, but we don't recommend it. How do the values $AICc$?

Prism also lets you choose to report the $AICc$. This would be useful only if you separately fit the same data to three or more models. You can then use the $AICc$ to choose between them. But note that it only makes sense to compare $AICc$ between fits, when the only difference is the model you chose. If the data or weighting are not identical between fits, then any comparison of $AICc$ values would be meaningless.

Residual tests. Are the residuals Gaussian with appropriate weighting?

Least-squares nonlinear regression assumes that the distribution of residuals follows a Gaussian distribution (robust nonlinear regression does not make this assumption). Prism can test this assumption by running a normality test on the residuals. Prism offers three normality tests. We recommend the D'Agostino-Pearson test.

Nonlinear regression assumes that, on average, the distance of the points from the curve is the same all the way along the curve, or that you have accounted for systematic differences by choosing an appropriate weighting. Prism can test this assumption with a test for appropriate weighting. If you have chosen equal weighting (the default) this is the same as a test for homoscedasticity.
Does the curve systematically deviate from the points?

Does the curve follow the trend of the data? Or does the curve systematically deviate from the trend of the data? Prism offers two tests that answer these questions.

If you have entered replicate Y values, choose the replicates test to find out if the points are 'too far' from the curve (compared to the scatter among replicates). If the P value is small, conclude that the curve does not come close enough to the data.

The runs test is available if you entered single Y values (no replicates) or chose to fit only the means rather than individual replicates (weighting tab). A 'run' is a series of consecutive points on the same side of the curve. If there are too few runs, it means the curve is not following the trend of the data. If you fit several curves at once, sharing one or more parameters (but not all the parameters) with global regression, Prism will report the runs test for each curve fit, but not for the global fit. Prior versions of Prism reported the overall runs test for the global fit, by summing the runs for each component curve, but this is not standard and Prism 7 does not report this.

If you choose a residual plot, Prism creates a new graph. The X axis is the same as the graph of the data, while the Y axis plots the distance of each point from the curve (the residuals). Points with positive residuals are above the curve; points with negative residuals are below the curve. Viewing a residual plot can help you assess whether the distribution of residuals is random above and below the curve.

Are the parameters intertwined or redundant or skewed?

What does it mean for parameters to be intertwined? After fitting a model, change the value of one parameter but leave the others alone. The curve moves away from the points. Now, try to bring the curve back so it is close to the points by changing the other parameter(s). If you can bring the curve closer to the points, the parameters are intertwined. If you can bring the curve back to its original position, then the parameters
are redundant. In this case, Prism will alert you by labeling the fit 'ambiguous'.

We suggest that you report the dependency and not bother with the covariance matrix. When you are getting started with curve fitting, it is OK to leave both options unchecked.

Even though nonlinear regression, as its name implies, is designed to fit nonlinear models, some of the inferences actually assume that some aspects of the model are close to linear, so that the distribution of each parameter is symmetrical. This means that if you analyzed many data sets sampled from the same system, the distribution of the best-fit values of the parameter would be symmetrical and Gaussian.

If the distribution of a parameter is highly skewed, the reported SE and CI for that parameter will not be very useful ways of assessing precision. Hougaard's measure of skewness quantifies how skewed each parameter is.

Could outliers impact the results?

Nonlinear regression is based on the assumption that the scatter of data around the ideal curve follows a Gaussian distribution. The presence of one or a few outliers (points much further from the curve than the rest) can overwhelm the least-squares calculations and lead to misleading results.

Check this option to count the outliers, but leave them in the calculations. Choose how aggressively to define outliers by adjusting the ROUT coefficient.

If you chose the option in the Fit tab to exclude outliers from the calculations, then this option to simply count outliers (in the Diagnostics tab) is not available.

Would it help to use stricter convergence criteria?

Would it help to use stricter convergence criteria?

Quick: Two iterations that change less than 0.01%

Automatically switch to strict convergence when needed
Nonlinear regression is an iterative process. It starts with initial values of the parameters, and then repeatedly changes those values to increase the goodness-of-fit. Regression stops when changing the values of the parameters makes a trivial change in the goodness of fit.

Prism lets you define the convergence criteria in three ways.

- **Quick.** If you are fitting huge data sets, you can speed up the fit by using the 'quick' definition of convergence. With this choice, nonlinear regression is defined to converge when two iterations in a row change the sum-of-squares by less than 0.01%.

- **Medium (default).** Nonlinear regression is defined to converge when five iterations in a row change the sum-of-squares by less than 0.0001%.

- **Strict.** If you are having trouble getting a reasonable fit, you might want to try the stricter definition of convergence. With this choice, the nonlinear regression iterations don't stop until five iterations in a row change the sum-of-squares by less than 0.0000001%. It won't help very often, but is worth a try. The only reason not to always use the strictest choice is that it takes longer for the calculations to complete. That won't matter with small data sets, but will matter with large data sets or when you run scripts to analyze many data tables.

### 3.4.3.10 Flags tab

This feature is useful if you analyze many sets of data (perhaps by running a script), and need a way to automatically flag poor fits that need to be inspected more carefully.

In the Flag tab, define the attributes that define poor fits you wish to flag. You can choose to flag a fit with any or all of the following attributes: low $R^2$, too few data points, too high a dependency for any parameter, a failed normality test of residuals, a failed runs or residual test, or too many outliers.
Some options may be gray if the corresponding option is gray in the Diagnostics tab. You can't flag, for example, fits with high dependency unless you chose the option to compute dependency.

What choices should you make? It depends on the kind of work you are doing. In some systems, you would only flag fits with $R^2$ less than 0.50. In other systems, you expect tight data and so would flag any fit with $R^2$ less than 0.90. Once you have fit a model to dozens or hundreds of similar data sets, you'll have a feel for what to expect, and when to flag dubious fits. Until then, don't flag fits.

What happens if a fit is flagged? Not much! The word "Flagged" appears at the top of the fit as shown below. Use this as a cue to take a more careful look at this data set and its fit.

When you run a Monte-Carlo simulation, there is a new option to ignore (not tabulate) results from fits that are flagged.

### 3.4.4 Graphing tips: Nonlinear regression

Prism automatically graphs the best-fit curves from nonlinear regression, and you'll rarely need to think about this step.

- **Tips** on graphing curves
• **Graphing prediction and confidence bands**

• **Adding an equation**, or a table of best-fit parameters, to the graph.

• **Graphing outliers**.

• **Graphing residuals**.

### 3.4.4.1 Graphing best-fit curves

**Automatic vs. manual plotting**

When Prism performs nonlinear regression, it automatically superimposes the curves on the graph.

If you need to create additional graphs, or change which curve is plotted on which graph, keep in mind that from Prism's point of view, a curve generated by nonlinear regression is simply a data set. You can add curves to a graph or remove curves from a graph on the 'Data on graph' tab of the Format Graph dialog.

**Plotting curves after comparing models**

If you ask Prism to compare two models, only the selected model is graphed. If you want to graph both models, you will need to analyze the data twice, once with each model.

**Plotting on logarithmic axes**

With Prism 5 and 6, it is Ok to stretch an axis to a log scale, a choice on top of the Format Axis dialog. With prior versions, this didn't work well. The curve was defined as a series of line segments, and these would be far from equally spaced on a log axis. Prism 5 and 6 are smarter, so the curve will look smooth on either a linear or logarithmic axis.

**Plotting only a portion of the curve**

Let's say that the X values of your data range from 0 to 100, and you fit a curve with nonlinear regression. Now you want to create a graph...
showing only the data with X values between 0 and 10. The curve will look ragged. The problem is that Prism defines the entire curve using 150 line segments. In this example, you only plot 10% of the range, so the curve will have only 15 line segments.

To fix this problem, go to the Range tab of nonlinear regression and choose to define the curve with more line segments. In this example, increasing the value tenfold, to 1500, would be about right.

3.4.4.2 Graphing confidence and prediction bands

The difference between confidence and prediction bands

The 95% confidence bands enclose the area that you can be 95% sure contains the true curve. It gives you a visual sense of how well your data define the best-fit curve.

The 95% prediction bands enclose the area that you expect to enclose 95% of future data points. This includes both the uncertainty in the true position of the curve (enclosed by the confidence bands), and also accounts for scatter of data around the curve.

Choose in the Diagnostics tab of the Nonlinear regression parameters dialog. Learn how they are calculated.

Prediction bands are always wider than confidence bands. When you have lots of data points, the discrepancy is huge.
Plotting confidence or prediction bands

After you choose to create a confidence or prediction band in the diagnostics tab, it should just appear on the graph. You can fine tune its appearance on the graph. Note the following:

- The fit curve is seen by the Format Graph dialog as one more data set to plot. Be sure to select one of those nonlinear fits first (top of dialog) before adjusting its properties.

- To plot the confidence bands, choose to show error bars, and set the style to ------ or ______.

- You can also choose to show an area fill within the error bands.

Why do so many points lie outside the confidence bands?

Confidence bands show you how well you know the location of the best fit line or curve. Given all the assumptions of the analysis, you can be 95% sure that the true curve (nonlinear regression) or line (linear regression) lies within the bands.
Prediction bands show you where you can expect the data to lie. You expect 95% of all data points to lie within the prediction bands.

With many data points, you expect a large fraction of the data points to lie outside the confidence band, but 95% to lie within the prediction bands. The confidence bands aren't supposed to show you the scatter of the data, but rather the uncertainty in the position of the line or curve. With lots of data, the line or curve is known fairly precisely, so only a small fraction of data lie within the bands. The figure below demonstrates this with linear regression.

Plotting both confidence and prediction bands

Prism lets you choose either a confidence band or a prediction band as part on the diagnostics tab. But not both. To plot both on one graph, analyze your data twice, choosing a confidence band the first time and a prediction band the second time. The curves are data sets that you can add to any graph by drag-n-drop or using the Change menu from the graph.

One-sided confidence (or prediction) bands

Confidence and prediction bands are usually two-sided. But you can get Prism to show the band on only one side.

One use for a one-sided confidence band would be on a graph that plots purity of a substance over time. You want to know, with a certain confidence, how low the purity could be at a certain time. You don't really care about quantifying how high the purity could be.
To plot a one-sided 95% confidence (or prediction) band:

1. Choose "90%" confidence bands in the Diagnostics tab of the nonlinear regression dialog. With 90% bands, there is a 5% chance that the true value is larger than the upper band, and 5% it is lower than the lower band. When you plot only one of these bands, therefore, it is really a 95% confidence band, which is probably what you want.

2. On the Format Symbols dialog, choose the best-fit line or curve and make sure that error bars are turned on with the "---" style. Then choose either Up or Down (but not Both) in the Direction drop-down menu.

**Viewing the XY coordinates of confidence or prediction bands**

1. Go to the Diagnostics tab and check the option to plot confidence or prediction bands.

2. Go to the Range tab of the nonlinear regression dialog.

3. Check the option to create a table of XY coordinates. This table will include the confidence or prediction bands.

4. When viewing the results, expand the results sheet in the navigator and go to the page labeled 'Curve'. For each X value, you'll see the y value and the distance the confidence or prediction curve is above and below that value.

**Situations where Prism can't plot confidence or prediction bands**

Prism will not plot confidence or prediction bands in several situations:

- If the best-fit value of a parameter hit a constraint, the fit is unlikely to be useful. Prism does not plot confidence or prediction bands, because they would almost certainly be misleading.

- If the results of nonlinear regression are ambiguous, the confidence or prediction bands would be super wide, maybe infinitely wide. They would not be useful, so Prism does not plot them.

- If you choose robust nonlinear regression, Prism does not compute confidence or prediction bands, as it cannot compute standard errors or confidence intervals of the parameters.
• The fit is perfect. If the sum-of-squares is 0.0 and R2 is 1.0, it is not possible to compute or interpret confidence or prediction bands.

• If the fit is interrupted, confidence and prediction bands are not computed.

• If you chose to weight the fit by 1/SD². To plot confidence and prediction bands, Prism needs to know the weighting factor at every X value in the range it plots the curve. Not a problem with most weighting schemes. But if you weight by the values you enter in the SD column, then Prism only knows those weights, and not the weights for X values between the points, so can't plot confidence or prediction bands.

3.4.4.3 Adding the equation to the graph

Prism does not automatically write equations on the graph. There are two approaches you can use to do so.

Microsoft Office comes with an equation editor. If Microsoft Equation Editor is installed on your system, you will see a quick-access button on your toolbar when you are on either a graph or a layout sheet.

You can create either a generic equation, or an equation with the best-fit values of the parameters substituted for the variable names.

If the button doesn't appear on your toolbar, create the equation inside of Microsoft Word or some other program, and then copy and paste.

Another approach is to simply show the parameter values from the curve of best fit in tabular form. Copy selected results and then paste onto a graph. The table will be linked, so its values will change if you edit or replace the data. This doesn't show the form of the equation (exponential decay, in this example) but gives you the results of this particular experiment, which may be more important.
The example below shows use of the equation editor to write both a generic and specific equation (for the control data set) and an embedded table showing the results for both data sets.

\[ Y = \text{Plateau} + (Y_0 - \text{Plateau})e^{-kX} \]
\[ Y = 987 + (9992 - 987)e^{-0.08927X} \]

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>Treated</th>
</tr>
</thead>
<tbody>
<tr>
<td>Y0</td>
<td>9992</td>
<td>9593</td>
</tr>
<tr>
<td>Plateau</td>
<td>987.0</td>
<td>1154</td>
</tr>
<tr>
<td>k</td>
<td>0.08927</td>
<td>0.3042</td>
</tr>
<tr>
<td>Half Life</td>
<td>7.765</td>
<td>2.278</td>
</tr>
<tr>
<td>Tau</td>
<td>11.20</td>
<td>3.287</td>
</tr>
<tr>
<td>Span</td>
<td>9005</td>
<td>8440</td>
</tr>
</tbody>
</table>

3.4.4.4 Graphing outliers

**How to identify or eliminate outliers**

Prism can identify outliers from the curve fit, and you can choose whether these should simply be plotted or also ignored by the curve fitting process. Choose to count the outliers on the Diagnostics tab, and to eliminate outliers on the Fit tab of the nonlinear regression dialog.

You can adjust the value of Q to redefine how aggressively Prism defines outliers.

**How Prism plots outliers**

If you choose to either count or eliminate outliers, Prism tabulates any outliers in a page of the results sheet called 'outliers' and plots them in red (unless data are plotted in red, in which case the outliers are blue).
The example above shows a graph where each replicate is plotted individually. The example below shows a graph which plots mean and SD error bars. The error bars are computed from all the data, including any outliers. The outliers are then superimposed on the graph as well.

Note that Prism does not remove the outlier from the regular data set. When you plot mean and error bar, the outlier is included in the calculation. When you plot individual data points, the outlier is still plotted with the full dataset. Then Prism superimposes the outlier(s) as a separate dataset plotted in front.

Look back at the first example, showing each replicate. You can see that the curve goes over the data points. If you want the curve to go under the data points, click the Reverse/Flip/Rotate button.
The results, above, are not exactly what you want. Indeed the relative front-to-back order of data and curve has reversed, so the curve is now behind the points. But the relative front-to-back order of the full dataset and the outlier dataset is also reversed, so the outlier is now behind the regular data point, so is invisible. This demonstrates that Prism does not really change the color of the data points of outliers. Rather, it plots the outliers twice -- once as part of the full data set and again as part of the outlier dataset superimposed on the graph. To get the effect you want here, go to the Data on Graph tab of the Format Graph dialog, and fine-tune the back-to-front order of data sets (with the outlier(s) and curve considered to be data sets).

3.4.4.5 Residual plot

When to plot residuals
A residual is the distance of a point from the curve. A residual is positive when the point is above the curve, and is negative when the point is below the curve. The residual table has the same X values as the original data, but the Y values are the vertical distances of the point from the curve.

Create a residual plot if you aren't sure that your data really follow the model you selected. Mild deviations of data from a model are often easier to spot on a residual plot.

**How to graph residuals**

Choose to create a residual plot by checking an option on the Diagnostics tab of the nonlinear regression dialog. Prism will automatically make a new graph. The residuals are tabulated in a separate page of results, and you can use these like any other table (make additional graphs, plot on other graphs, transform...).

**Interpreting a residual plot**

The X axis of the residual plot is the same as the graph of the data, while the Y axis is the distance of each point from the curve. Points with positive residuals are above the curve; points with negative residuals are below the curve.

An example is shown below, with a graph of the data and curve combined with a residual plot in a layout. If you look carefully at the curve on the left, you will see that the data points are not randomly distributed above and below the curve. There are clusters of points at early and late times that are below the curve, and a cluster of points at middle time points that are above the curve. This is much easier to see on the graph of the residuals in the inset. The data are not randomly scattered above and below the X-axis.
Residuals with weighted fits

If you choose (or accept the default) standard weighting, then the residuals are the difference between the actual Y value you entered and the Y value predicted by the model. If the data point is above the curve, the residual is positive. If the data point is below the curve, the residual is negative. Least-squares regression works to minimize the sum of the squares of these residuals.

If you choose another weighting scheme, Prism adjusts the definition of the residuals accordingly. The residual that Prism tabulates and plots equals the residual defined in the prior paragraph, divided by the weighting factor. The most common alternative weighting is "Weight by $1/Y^2$ (minimize relative distances squared)". In this case, the residual is defined to be the distance of the point from the curve divided by the Y value of the curve. Weighted nonlinear regression minimizes the sum of the squares of these residuals.

Note the ambiguity in defining weighting. The Prism dialog gives the choice to weight by $1/Y^2$. This means that the squared residual is divided by $Y^2$. The weighted residual is defined as the residual divided by Y. Prism minimizes the sum of the squares of these weighted residuals.

Earlier versions of Prism (up to Prism 4) always plotted basic unweighted residuals, even if you chose to weight the points unequally.

3.5 Interpreting nonlinear regression results

Interpreting nonlinear regression results can be confusing.
• If Prism reported error message on the top of the results column, here are the explanations.

• Help with graphing best-fit curves, with their confidence or prediction bands and residual plots.

• Read detailed explanations of the results, and review analysis checklists. Where to go depends on your goal.

Goal: Determine best-fit values
Analysis checklist
Learn more.

Goal: Compare models
Analysis checklist
Learn more.

Goal: Interpolate unknowns from a standard curve
Analysis checklist
Learn more.

3.5.1 Interpreting results: Nonlinear regression

When evaluating nonlinear regression results, first consider what your goal is. If your goal is to interpolate unknowns from a standard curve, skip this page and go right to this analysis checklist.

If your goal is to determine the values of the best-fit parameters or to compare models, answer the three questions below before looking at parameter values, R2, etc.

• Are your results free of error messages? Prism sometimes puts a short message at the top of the table of results. Read about the meaning of "Bad initial values", "Interrupted", "Not converged", "Ambiguous", "Hit constraint", "Don't fit", "Too few points", "Perfect fit", "Impossible weights", and "Equation not defined".
• Does the curve go near your data points? In rare cases, the curve may be far from the data points. This may happen, for example, if you picked the wrong equation. Look at the graph to make sure this didn't happen.

• Are the best-fit values of the parameters scientifically sensible? Nonlinear regression programs have no common sense and don't know the context of your experiment. The curve fitting procedure can sometimes yield results that make no scientific sense. For example with noisy or incomplete data, nonlinear regression can report a best-fit rate constant that is negative, a best-fit fraction that is greater than 1.0, or a best-fit Kd value that is negative. All these results are scientifically meaningless. Also check whether the best-fit values of the variables make sense in light of the range of the data. The results make no sense if the top plateau of a sigmoid curve is far larger than the highest data point, or an EC50 is not within the range of your X values. If the results make no scientific sense, they are unacceptable, even if the curve comes close to the points and R2 is close to 1.0.

If the answer to all three questions is yes, then it makes sense to delve into the numerical results of nonlinear regression in detail.

- **Standard errors and confidence intervals of parameters**
- **Normality tests of residuals**
- **Goodness of fit of nonlinear regression**
- **Runs test**
- **Replicates test**
- **Dependency and covariance matrix**
- **Hougaard's measure of skewness**
- **Could the fit be a local minimum?**
- **Outliers**
3.5.1.1 Best-fit parameter values

Before trying to interpret the rest of the results, first look at the best-fit value of each parameter in your model. First make sure you know what units each parameter is expressed in (if there are any units; some parameters are unitless constants). Next, make sure each value makes biological or scientific sense. If a value doesn't make any sense, ask your self these questions:

- Does your data actually define all the parameters?
- Should you be constraining a parameter (or several) to constant values? If not, should you be constraining them to have a range of values (only positive numbers, perhaps)?
- Should you be fitting a family of datasets together using global nonlinear regression?

3.5.1.2 Standard error of parameters

Interpreting the standard errors of parameters

The only real purpose of the standard errors is as an intermediate value used to compute the confidence intervals. If you want to compare Prism's results to those of other programs, you will want to include standard errors in the output. Otherwise, we suggest that you ask Prism to report the confidence intervals only (choose on the Diagnostics tab). The calculation of the standard errors depends on the sum-of-squares, the spacing of X values, the choice of equation, and the number of replicates.

'Standard error' or 'standard deviation'? 

Prism reports the standard error of each parameter, but some other programs report the same values as 'standard deviations'. Both terms mean the same thing in this context.

When you look at a group of numbers, the standard deviation (SD) and standard error of the mean (SEM) are very different. The SD tells you about the scatter of the data. The SEM tells you about how well you have determined the mean. The SEM can be thought of as "the standard deviation of the mean" -- if you were to repeat the experiment many
times, the SEM (of your first experiment) is your best guess for the standard deviation of all the measured means that would result.

When applied to a calculated value, the terms "standard error" and "standard deviation" really mean the same thing. The standard error of a parameter is the expected value of the standard deviation of that parameter if you repeated the experiment many times. Prism (and most programs) calls that value a standard error, but some others call it a standard deviation.

### 3.5.1.3 Confidence intervals of parameters

**Do not ignore the confidence intervals**

In most cases, the entire point of nonlinear regression is to determine the best-fit values of the parameters in the model. The confidence interval tells you how tightly you have determined these values. If a confidence interval is very wide, your data don't define that parameter very well. Confidence intervals are computed from the standard errors of the parameters.

**How accurate are the standard errors and confidence intervals?**

Prism gives you two choices in the Confidence tab.

The asymptotic or approximate standard errors reported by Prism (and virtually all other nonlinear regression programs) are based on some mathematical simplifications. They are calculated assuming that the equation is linear, but are applied to nonlinear equations. This simplification means that the intervals can be too optimistic.

Prism also lets you choose to report asymmetrical confidence intervals (profile likelihood intervals) and these are more accurate.

**Sometimes Prism reports "very wide" instead of reporting the confidence interval**

If you see the phrase 'very wide' instead of a confidence interval, you will also see the phrase 'ambiguous' at the top of the results tables. This means that the data do not unambiguously define the parameters. Many sets of parameters generate curves that fit the data equally well. The curve may fit well, making it useful artistically or to interpolate unknowns, but you can't rely on the best-fit parameter values.
Confidence intervals of transformed parameters

In addition to reporting the confidence intervals of each parameter in the model, Prism can also report confidence intervals for transforms of those parameters. For example, when you fit an exponential model to determine the rate constant, Prism also fits the time constant tau, which is the reciprocal of the rate constant.

When you write your own equation, or clone an existing one, choose between two ways to compute the confidence interval of each transformed parameter. If you pick a built-in equation, Prism always reports asymmetrical confidence intervals of transformed parameters.

Do not mix up confidence intervals and confidence bands

It is easy to mix up confidence intervals and confidence bands. Choose both on the Diagnostics tab.

The 95% confidence interval tells you how precisely Prism has found the best-fit value of a particular parameter. It is a range of values, centered on the best-fit value. Prism can display this range in two formats:

The 95% confidence bands enclose the area that you can be 95% sure contains the true curve. It gives you a visual sense of how well your data define the best-fit curve. It is closely related to the 95% prediction bands, which enclose the area that you expect to enclose 95% of future data points. This includes both the uncertainty in the true position of the curve (enclosed by the confidence bands), and also accounts for scatter of data around the curve. Therefore, prediction bands are always wider than confidence bands.
3.5.1.4 Normality tests of residuals

Interpreting a normality test

The result of a normality test is expressed as a P value that answers this question:

If your model is correct and all scatter around the model follows a Gaussian population, what is the probability of obtaining data whose residuals deviate from a Gaussian distribution as much (or more so) as your data does?

If the P value is large, then the residuals pass the normality test. If the P value is small, the residuals fail the normality test and you have evidence that your data don't follow one of the assumptions of the regression. Things to consider:

- Fit a different model
- Weight the data differently.
- Exclude outliers.

A large P value means that your data are consistent with the assumptions of regression (but certainly does not prove that the model is correct). With small numbers of data points, normality tests have little power to detect modest deviations from a Gaussian distribution.
How the normality tests work

We recommend relying on the D’Agostino-Pearson normality test. It first computes the skewness and kurtosis to quantify how far from Gaussian the distribution is in terms of asymmetry and shape. It then calculates how far each of these values differs from the value expected with a Gaussian distribution, and computes a single P value from the sum of these discrepancies. It is a versatile and powerful (compared to some others) normality test, and is recommended. Note that D’Agostino developed several normality tests. The one used by Prism is the "omnibus K2" test.

An alternative is the Shapiro-Wilk normality test. We prefer the D'Agostino-Pearson test for two reasons. One reason is that, while the Shapiro-Wilk test works very well if every residual is unique, it does not work well when several residuals are identical. The other reason is that the basis of the test is hard for non mathematicians to understand.

Earlier versions of Prism offered only the Kolmogorov-Smirnov test. We still offer this test (for consistency) but no longer recommend it. This test compares the cumulative distribution of the data with the expected cumulative Gaussian distribution, and bases its P value simply on the largest discrepancy. This is not a very sensitive way to assess normality, and we now agree with this statement: "The Kolmogorov-Smirnov test is only a historical curiosity. It should never be used."

The Kolmogorov-Smirnov method as originally published assumes that you know the mean and SD of the overall population (perhaps from prior work). When analyzing data, you rarely know the overall population mean and SD. You only know the mean and SD of your sample. To compute the P value, therefore, Prism uses the Dallal and Wilkinson approximation to Lilliefors’ method (Am. Statistician, 40:294-296, 1986). Since that method is only accurate with small P values, Prism simply reports “P>0.10” for large P values.

Reference

3.5.1.5  R squared

**Meaning of R²**

Key points about R²

- The value R² quantifies goodness of fit.
- It is a fraction between 0.0 and 1.0, and has no units. Higher values indicate that the model fits the data better.
- When R² equals 0.0, the best-fit curve fits the data no better than a horizontal line going through the mean of all Y values. In this case, knowing X does not help you predict Y.
- When R²=1.0, all points lie exactly on the curve with no scatter. If you know X you can calculate Y exactly.
- You can think of R² as the fraction of the total variance of Y that is explained by the model (equation). With experimental data (and a sensible model) you will always obtain results between 0.0 and 1.0.
- There is really no general rule of thumb about what values of R² are high, adequate or low. If you repeat an experiment many times, you will know what values of R² to expect, and can investigate further when R² is much lower (or higher) than the expected value.
- By tradition, statisticians use uppercase (R²) for the results of nonlinear and multiple regression and lowercase (r²) for the results of linear regression, but this is a distinction without a difference.
- In linear regression, the R² compares the fits of the best fit regression line with a horizontal line (forcing the slope to be 0.0). The horizontal line is the simplest case of a regression line, so this makes sense. With many models used in nonlinear regression, the horizontal line is not a simple case and can't be generated at all from the model. So comparing the fits of the chosen model with the fit of a horizontal line doesn't quite make sense. For this reason Minitab does not report R² with nonlinear regression and SAS calls the value "Pseudo R²".
- Statisticians have actually developed eight different methods for computing R² for nonlinear regression(1)! As detailed below, Prism uses method 1 from Kvalseth's paper.
Don't overemphasize $R^2$

A common mistake is to use $R^2$ as the main criteria for whether a fit is reasonable. A high $R^2$ tells you that the curve came very close to the points. That doesn't mean the fit is "good" in other ways. The best-fit values of the parameters may have values that make no sense (for example, negative rate constants) or the confidence intervals may be very wide. The fit may be ambiguous. You need to look at all the results to evaluate a fit, not just the $R^2$.

How $R^2$ is calculated -- unweighted fits

$R^2$ is computed from the sum of the squares of the distances of the points from the best-fit curve determined by nonlinear regression. This sum-of-squares value is called SSreg, which is in the units of the Y-axis squared. To turn $R^2$ into a fraction, the results are normalized to the sum of the square of the distances of the points from a horizontal line through the mean of all Y values. This value is called SSTot. If the curve fits the data well, SSres will be much smaller than SSTot.

The figure below illustrates the calculation of $R^2$. Both panels show the same data and best-fit curve. The left panel also shows a horizontal line at the mean of all Y values, and vertical lines showing how far each point is from the mean of all Y values. The sum of the square of these distances (SSTot) equals 62735. The right panel shows the vertical distance of each point from the best-fit curve. The sum of squares of these distances (SSres) equals 4165.

$$R^2 = 1.0 - \frac{SSres}{SSTot} = 1.0 - \frac{4165}{62735} = 0.9336$$

If you share parameters (perform global nonlinear regression) SSRes in the equation above is the Sum of Squares reported by Prism in Global result column for model with shared parameters being fitted, and SSTot is the Sum of Squares of each Y value (from each data set) around the mean of ALL Y values (from all data sets).
How $R^2$ is calculated -- weighted fits

There does not seem to be an established method for computing the $R^2$ of a weighted nonlinear fit. Prism (since 6.00) calculates as detailed below. But note that earlier versions of Prism computed the $R^2$ of weighted fits differently.

1. Fit the model using relative ($1/Y^2$) or Poisson ($1/Y$) or general ($1/X^K$) weighting to compute the weighted sum-of-squares (wSSmodel). This is the weighted sum-of-squares of the residuals from your model.

2. Remember the weight each point received.

3. Fit the data to a horizontal line model ($Y=\text{Mean} + 0*X$) using the same weights for each point as used in step 1. This is the weighted sum-of-squares of the residuals from a horizontal line (wSShorizontal).

The weighted $R^2$ is:

$$1.0 - \frac{\text{wSSmodel}}{\text{wSShorizontal}}$$

Note that weights are only computed when fitting the model. Prism does not compute weights for the fit of the horizontal line but rather uses exactly the same weights as were used to fit the model. This ensures that the sum of the weights is identical in both fits.

$R^2$ can be negative!

Appearances can be deceptive. $R^2$ is not really the square of anything. If SSres is larger than SStot, $R^2$ will be negative (see equation above). While it is surprising to see something called "squared" have a negative value, it is not impossible (since $R^2$ is not actually the square of R).

How can this happen? SSres is the sum of the squares of the vertical distances of the points from the best-fit curve (or line). SStot is the sum of the squares of the vertical distances of the points from a horizontal line drawn at the mean Y value. SSres will exceed SStot when the best-fit line or curve fits the data even worse than does a horizontal line.

$R^2$ will be negative when the best-fit line or curve does an awful job of fitting the data. This can only happen when you fit a poorly chosen model (perhaps by mistake), or you apply constraints to the model that don't make any sense (perhaps you entered a positive number when you
intended to enter a negative number). For example, if you constrain the Hill slope of a dose-response curve to be greater than 1.0, but the curve actually goes downhill (so the Hill slope is negative), you might end up with a negative $R^2$ value and nonsense values for the parameters.

Below is a simple example. The blue line is the fit of a straight line constrained to intercept the Y axis at $Y=150$ when $X=0$. SSres is the sum of the squares of the distances of the red points from this blue line. SSTot is the sum of the squares of the distances of the red points from the green horizontal line. Since Sres is much larger than SSTot, the $R^2$ (for the fit of the blue line) is negative.

$$R^2 = 1 - \frac{SS_{res}}{SS_{tot}} = 1 - \frac{11623}{1320} = -7.0$$

Prism file

If $R^2$ is negative, check that you picked an appropriate model, and set any constraints correctly.

Adjusted $R^2$

The $R^2$ quantifies how well a model fits the data. When you compare models, the one with more parameters can bend and twist more to come nearer the points, and so almost always has a higher $R^2$. This is a bit misleading.

The adjusted $R^2$ accounts for the number of parameters fit by the regression, and so can be compared between models with different numbers of parameters. The equations for the regular and adjusted $R^2$ are compared here (SSresiduals is the sum-of-squares of the discrepancy
between the Y value of the curve and the data; SS\text{total} is the sum-of-squares of the differences between the overall Y mean and each Y value; n is the number of data points, and K is the number of parameters fit):

\[ R^2 = 1 - \frac{SS_{\text{residuals}}}{SS_{\text{total}}} \]

\[
\text{Adjusted } R^2 = 1 - \frac{SS_{\text{residuals}}}{SS_{\text{total}} (n - 1)} \]

The adjusted \( R^2 \) is smaller than the ordinary \( R^2 \) whenever the number of parameters (K) is greater than 1.


3.5.1.6 Sum-of-squares

Nonlinear regression finds the curve that minimizes the sum of square of the distances of the points from the curve. So Prism reports that sum-of-square value. This is useful if you want to compare Prism with another program, or compare two fits manually. Otherwise, the value is not very helpful.

If you choose to differentially weight your data, Prism reports both the absolute and the weighted sum-of-squares.

3.5.1.7 Standard deviation of the residuals

Prism can quantify goodness of fit by reporting the standard deviation of the residuals, computed in three distinct ways. Remember that the residual is the vertical distance (in Y units) of the point from the fit line
or curve. If you have n data points, after the regression, you have n residuals.

**What units? How to interpret?**

All three values (RMSE, Sy.x, and RSDR) are expressed in the same units as Y and all can be interpreted in roughly the same way as the typical deviation of the points from the line or curve.

**RMSE**

If you simply take the standard deviation of those n values, the value is called the root mean square error, RMSE. The mean of the residuals is always zero, so to compute the SD, add up the sum of the squared residuals, divide by n-1, and take the square root:

\[
RMSE = \sqrt{\frac{\sum(residual^2)}{n - 1}}
\]

Prism will report the RMSE when you check the appropriate option in the Diagnostics tab, because some fields use it. But we recommend reporting the Sy.x (below).

**Sy.x or Se**

The Sy.x is computed in a very similar way, but the denominator is n-K, where K is the number of parameters fit by regression. The value n-K is the number of degrees of freedom of the regression. If you only fit one parameter, then the RMSE and Sy.x are the same. If you fit two or more parameters, the Sy.x is larger and is a better estimate of goodness-of-fit. We recommend reporting it rather than the RMSE.

\[
Sy.x = \sqrt{\frac{\sum(residual^2)}{n - K}}
\]
RSDR

If you chose robust regression, Prism computes a different value we call the Robust Standard Deviation of the Residuals (RSDR). The goal here is to compute a robust standard deviation, without being influenced by outliers. In a Gaussian distribution, 68.27% of values lie within one standard deviation of the mean. We therefore calculate this value, which we call P68. It turns out that this value underestimates the SD a bit, so the RSDR is computed by multiplying the P68 by \( n/(n-K) \), where K is the number of parameters fit.

3.5.1.8 Why Prism doesn't report the chi-square of the fit

Some programs report a chi-square value with the results of nonlinear regression. Prism doesn't. This page explains how the chi-square value could be useful and why we don't report it.

Is the sum-of-squares too high?

Nonlinear regression minimizes the sum of the squared vertical distances between the data point and the curve. But how can you interpret the sum-of-squares, abbreviated SS? You can't really, as it depends on the number of data points you collected and the units you used to express Y.

The chi-square approach is to compare the observed scatter of the points around the curve (SS) with the amount of experimental scatter you expect to see based theory. This is done by computing chi-square using this equation:

\[
\chi^2 = \sum_{i=1}^{i=n} \left( \frac{Y_i - Y_{\text{curve}}}{\sigma_i} \right)^2
\]

Chi-square is the sum of the square of the ratio of the distance of a point from the curve divided by the predicted standard deviation at that value of X. Note that the denominator is the predicted standard deviation, not the actual standard deviation computed in this particular experiment.
If you know that the SD is the same for all values of X, this simplifies to:

\[ \chi^2 = \frac{\sum_{i=1}^{n} (Y_i - Y_{\text{curve}})^2}{SD^2} \]

The standard deviation value must be computed from lots of data so the SD is very accurate. Or, better, the SD can come from theory.

If you assume that replicates are scattered according to a Gaussian distribution with the SD you entered, and that you fit the data to the correct model, then the value of chi-square computed from that equation will follow a known chi-square distribution. This distribution depends on the number of degrees of freedom, which equals the number of data points minus the number of parameters. Knowing the value of chi-square and the number of degrees of freedom, a P value can be computed.

How can you interpret a small P value? If you are quite sure the scatter really is Gaussian, and that predicted SD is correct, a small P value tells you that your model is not right -- that the curve really doesn't follow the data very well. You should seek a better model.

But often a low P value just tells you that you don't predict the SD as well as you thought you would. It is hard to determine the SD values precisely, so hard to interpret the chi-square value. For this reason, Prism doesn't attempt the chi-square computation. We fear it would be more misleading than helpful.

**Alternatives**

Several approaches have been devised to answer the question of whether the SS is too high:

- The value of sum-of-squares can be used to compute $R^2$. This value is computed by comparing the sum-of-squares (a measure of scatter of points around the curve) with the total variation in Y values (ignoring X, ignoring the model). What values of $R^2$ do you expect? How low a value is too low? You can’t really answer that in general, as the answer depends on your experimental system.

- If you have collected replicate Y values at each value of X, you can compare the SS with a value predicted from the scatter among
replicates. Prism calls this the replicates test. This is very useful, but only if you have collected replicate Y measurements at each X.

- You can propose alternative models, and compare their fit to the data

**Summary**

Chi-square compares the actual discrepancies between the data and the curve with the expected discrepancies (assuming you selected the right model) based on the known SD among replicates. If the discrepancy is high, then you have some evidence that you've picked the wrong model. The advantage of the chi-square calculation is that it tests the appropriateness of a single model, without having to propose an alternative model and without having to have replicate values. The disadvantage is that the calculation depends on knowing the SD values with sufficient precision, which is often not the case.

Recommended alternatives are to compare the fits of two models, or use the replicates test.

3.5.1.9 Runs test

**Goal of the runs test**

The runs test asks whether the curve deviates systematically from your data. The runs test is useful only if you entered single Y values (no replicates) or chose to fit only the means rather than individual replicates (weighting tab). If you entered and analyzed replicate data, use the replicates test instead.

A run is a series of consecutive points that are either all above or all below the regression curve. Another way of saying this is that a run is a consecutive series of points whose residuals are either all positive or all negative. After fitting a curve, Prism counts the actual number of runs and calculates the predicted number of runs (based on number of data points). The runs test compares these two values.

**How the runs test works**

If the data points are randomly distributed above and below the regression curve, it is possible to calculate the expected number of runs.
If there are Na points above the curve and Nb points below the curve, the number of runs you expect to see equals \((2NaNb)/(Na+Nb)\)+1.

If the model fits the data poorly, you will tend to see clusters of points on the same side of the curve. This means you will have fewer runs than predicted from sample size, and the runs test will produce a low P value.

**Interpreting the P value from a runs test**

The P value answers this question:

If the data are randomly scattered above and below the curve, what is the probability of observing as few runs (or even fewer) than actually observed in this analysis?

If the runs test reports a low P value, conclude that the curve doesn't describe the data very well. The problem might be that some of the errors are not independent, that outliers are mucking up the fit, or that you picked the wrong model.

Note that the P value is one-tailed. If you observed more runs than expected, the P value will be higher than 0.50.

**Runs test with global fits**

Prism reports the runs test for each data set, but does not report a global runs test. Prior versions of Prism reported a global runs test, but it is not clear if that result was actually meaningful.

**3.5.1.10 Replicates test**

**Goal of the replicates test**

When evaluating a nonlinear fit, one question you might ask is whether the curve is 'too far' from the points. The answer, of course, is another question: Too far compared to what? If you have collected one Y value at each X value, you can't really answer that question (except by referring to other similar experiments). But if you have collected replicate Y values at each X, then you can ask whether the average distance of the points from the curve is 'too far' compared to the scatter among replicates.
If you have entered replicate Y values, choose the replicates test to find out if the points are 'too far' from the curve (compared to the scatter among replicates). If the P value is small, conclude that the curve does not come close enough to the data.

Example

![Graph of dose response](image)

The response at the last two doses dips down a bit. Is this coincidence? Or evidence of a biphasic response?

One way to approach this problem is to specify an alternative model, and then compare the sums-of-squares of the two fits. In this example, it may not be clear which biphasic model to use as the alternative model. And there probably is no point in doing serious investigation of a biphasic model in this example, without first collecting data at higher doses.

Since replicate values were obtained at each dose, the scatter among those replicates lets us assess whether the curve is ‘too far’ from the points.

After checking the option (on the Diagnostics tab) to perform the replicates test, Prism reports these results:
The value in the first row quantifies the scatter of replicates, essentially pooling the standard deviations at all X values into one value. This value is based only on variation among replicates. It can be computed without any curve fitting. If the replicates vary wildly, this will be a high value. If the replicates are all very consistent, this value will be low.

The value in the second row quantifies how close the curve comes to the mean of the replicates. If the curve comes close to the mean of the replicates, the value will be low. If the curve is far from some of those means, the value will be high.

The third row (F) is the square of the ratio of those two SD values.

If the model is correct, and all the scatter is Gaussian random variation around the model, then the two SD values will probably be similar, so F should be near 1. In this example, F is greater than 1 because the SD for lack of fit is so much greater than the SD of replicates. The P value answers this question:

If the model was chosen correctly and all scatter is Gaussian, what is the chance of finding a F ratio so much greater than 1.0?

A small P value is evidence that the data actually follow a model that is different than the model that you chose. In this example, this suggests that maybe some sort of biphasic dose-response model is correct -- that the dip of those last few points is not just coincidence.

How the replicates test calculations are done

This test for lack of fit is discussed in detail in advanced texts of linear regression (1,2) and briefly mentioned in texts of nonlinear regression (3, 4). Here is a brief explanation of the method:

The SD of replicates is computed by summing the square of the distance of each replicate from the mean of that set of replicates. Divide that sum
by its degrees of freedom (the total number of points minus the number of X values), and take the square root. This value is based only on variation among replicates. It can be computed without any curve fitting.

The SD lack of fit is a bit trickier to understand. Replace each replicate with the mean of its set of replicates. Then compute the sum of square of the distance of those points from the curve. If there are triplicate values, then each of the three replicate values is replaced by the mean of those three values, the distance between that mean and the curve is squared, and that square is entered into the sum three times (one for each of the replicates). Now divide that sum of squares by its degree of freedom (number of points minus number of parameters minus number of X values), and take the square root.

The F ratio is the square of the ratio of the two SDs.

The P value is computed from the F ratio and the two degree of freedom values, defined above.

References


3.5.1.11 Dependency of each parameter

**Intertwined parameters**

When your model has two or more parameters, as is almost always the case, the parameters can be intertwined.

What does it mean for parameters to be intertwined? After fitting a model, change the value of one parameter but leave the others alone. this will move the curve away from the points. Now change the other parameter(s) in an attempt to move the curve close to the data points. If you can bring the curve closer to the points, the parameters
are intertwined. If you can bring the curve back to its original position, then the parameters are completely redundant.

Prism can quantify the relationships between parameters in two ways. If you are in doubt, we suggest that you focus on the dependency values and not bother with the covariance matrix.

### Dependency

#### What is dependency and how do I ask Prism to compute it?

Dependency is reported for each parameter, and quantifies the degree to which that parameter is intertwined with others. Check a check box on the Diagnostics tab of nonlinear regression to view dependencies for each parameter.

#### Interpreting dependency

The value of dependency always ranges from 0.0 to 1.0.

A dependency of 0.0 is an ideal case when the parameters are entirely independent (mathematicians would say orthogonal). In this case the increase in sum-of-squares caused by changing the value of one parameter cannot be reduced at all by also changing the values of other parameters. This is a very rare case.

A dependency of 1.0 means the parameters are redundant. After changing the value of one parameter, you can change the values of other parameters to reconstruct exactly the same curve. If any dependency is greater than 0.9999, GraphPad labels the fit 'ambiguous'.

With experimental data, of course, the value will almost always lie between these extremes. Clearly a low dependency value is better. But how high is too high? Obviously, any rule-of-thumb is arbitrary. But dependency values up to 0.90 and even 0.95 are not uncommon, and are not really a sign that anything is wrong.

A dependency greater than 0.99 is really high, and suggests that something is wrong. This means that you can create essentially the same
curve, over the range of X values for which you collected data, with multiple sets of parameter values. Your data simply do not define all the parameters in your model. If your dependency is really high, ask yourself these questions:

- Can you fit to a simpler model?
- Would it help to collect data over a wider range of X, or at closer spaced X values? It depends on the meaning of the parameters.
- Can you collect data from two kinds of experiments, and fit the two data sets together using global fitting?
- Can you constrain one of the parameters to have a constant value based on results from another experiment?

If the dependency is high, and you are not sure why, look at the covariance matrix (see below). While the dependency is a single value for each parameter, the covariance matrix reports the normalized covariance for each pair of parameters. If the dependency is high, then the covariance with at least one other parameter will also be high. Figuring out which parameter that is may help you figure out where to collect more data, or how to set a constraint.

**How dependency is calculated**

This example will help you understand how Prism computes dependency.

### 3.5.1.12 Covariance matrix

**What is the covariance matrix and how do I ask Prism to compute it?**

The normalized covariance is reported for each pair of parameters, and quantifies the degree to which those two parameters are intertwined. Check a check box on the Diagnostics tab of nonlinear regression to view this covariance matrix.
Interpreting the normalized covariance matrix

Each value in the normalized covariance matrix ranges from -1.0 to 1.0. A value equal to -1.0 or 1.0 means the two parameters are redundant. A value of 0.0 means the parameters are completely independent or orthogonal -- if you change the value of one parameter you will make the fit worse and changing the value of the other parameter can't make it better. You can interpret a normalized covariance much as you interpret a correlation coefficient.

Note the difference between covariance and dependency. Each value in the covariance matrix tells you how much two parameters are intertwined. In contrast, each dependency value tells you how much that parameter is intertwined with all other parameters.

How to convert to the nonnormalized variance/covariance matrix

Some other programs report the actual (not normalized) variance-covariance matrix. Compute the actual covariance -- cov(i,j) -- of any two parameters (so i does not equal j) from the normalized matrix Prism reports -- NormCov(i,j) -- and the standard errors of the parameters using this equation:

\[ \text{Cov}(i, j) = \text{NormCov}(i, j) \times \text{SE}(i) \times \text{SE}(j) \]

Prism does not report the normalized covariance matrix for a parameter with itself, because the normalized covariance of any parameter with itself equals, by definition, 1.0. The covariance of any parameter with itself is better called its variance. You can calculate the variance of any parameter (a diagonal value in the variance-covariance matrix) using this equation:

\[ \text{Cov}(i, i) = \text{SE}(i)^2 \]

3.5.1.13 Confidence and prediction bands

What are confidence and prediction bands?
The 95% **confidence bands** enclose the area that you can be 95% sure contains the true curve. It gives you a visual sense of how well your data define the best-fit curve. It is closely related to the 95% **prediction bands**, which enclose the area that you expect to enclose 95% of future data points. This includes both the uncertainty in the true position of the curve (enclosed by the confidence bands), and also accounts for scatter of data around the curve. Therefore, prediction bands are always wider than confidence bands. **How confidence and prediction bands are computed.**

![Graph showing 95% confidence and prediction bands](image)

**How to plot confidence bands, prediction bands or both**

Choose to plot confidence or prediction bands by an checking option on the **Diagnostics tab**.

Prism lets you choose either a confidence band or a prediction band, but not both. To plot both on one graph, you need to analyze your data twice, choosing a confidence band the first time and a prediction band the second time.

The regression lines or curves are data sets that you can add to any graph by drag-n-drop or using the Change menu from the graph.

**How to view the XY coordinates of the confidence or prediction bands**

Prism can also show you the XY coordinates that define the curve and confidence bands (or prediction bands). But it only shows this table if you ask for it:

1. Go to the Diagnostics tab and check the option to plot confidence or prediction bands.
2. Go to the Range tab of the nonlinear regression dialog.

3. Check the option to create a table of XY coordinates. This table will include the confidence or prediction bands.

4. Enter the number of XY pairs you want to define the curve. 150 is a reasonable starting value, but of course it all depends on why you want these values.

5. When viewing the results, expand the results sheet in the navigator and go to the page labeled 'Curve'. For each X value, you'll see the Y value and the distance the confidence or prediction curve is above and below that value.

How to plot a one-sided confidence or prediction band

In some situations, it makes sense to plot only one side of the confidence or prediction band. One use for a one-sided confidence band would be on a graph that plots purity of a substance over time. You want to know, with a certain confidence, how low the purity could be at a certain time. But you don't really care about quantifying how high the purity could be.

To plot a one sided 95% confidence or prediction band:

1. Check the option (on the Diagnostics tab) to plot "90%" confidence (or prediction) bands. When you plot in only one direction, this is really a 95% confidence band.

2. On the Format Graph dialog, choose the data set that defines regression curve and make sure that error bars are turned on with the "---" style.

3. Choose to plot those error bands in one direction.

![Show error bars](dropdown)

- Color: [ ]
- Dir.: Above [ ]
- Style: [ ]
- Thickness: 1 pt [ ]

Do not mix up confidence intervals and confidence bands

It is easy to mix up confidence intervals and confidence bands. Choose both on the Diagnostics tab.
The **95% confidence interval** tells you how precisely Prism has found the best-fit value of a particular parameter. It is a range of values, centered on the best-fit value. Prism can display this range in two formats:

Don't mix up confidence bands and prediction bands

Note the difference between confidence and prediction bands:

The 95% confidence bands enclose the area that you can be 95% sure contains the true curve. If you have many data points, the confidence bands will be near the line or curve, and most of your data will lie outside the confidence bands.

The 95% prediction bands enclose the area that you expect to enclose 95% of future data points. They are wider than confidence bands -- much wider with large data sets.

Situations where Prism won't plot confidence or prediction bands

Prism will not plot confidence or prediction bands in several situations:

- If the best-fit value of a parameter **hits a constraint**, the fit is unlikely to be useful. Prism does not plot confidence or prediction bands, because they would almost certainly be misleading.

- If the results of nonlinear regression are **ambiguous**, the confidence or prediction bands would be super wide, maybe infinitely wide. They would not be useful, so Prism does not plot them.

- If you choose robust nonlinear regression, Prism does not compute confidence or prediction bands, as it cannot compute standard errors or confidence intervals of the parameters.

- The fit is **perfect**. If the sum-of-squares is 0.0 and R2 is 1.0, it is not possible to compute or interpret confidence or prediction bands.
• If the fit is interrupted, confidence and prediction bands are not computed.

• If you chose to weight by $1/\text{SD}^2$. To compute confidence or prediction bands, Prism needs to be able to compute the weights for any value of $X$. When you weight by $1/\text{SD}^2$, weights are only known at the $X$ values of the data, but not for other $X$ values, so Prism cannot compute prediction or confidence bands.

3.5.1.14 Hougaard's measure of skewness

Consequences of an asymmetrical parameter

Even though nonlinear regression, as its name implies, is designed to fit nonlinear models, some of the inferences actually assume that some aspects of the model are close to linear, so that the uncertainty about each parameter’s value is symmetrical. This means that if you analyzed many data sets sampled from the same system, the distribution of the best-fit values of the parameter would be symmetrical and Gaussian.

If the distribution of a parameter is highly skewed, there are two consequences:

• The SE of that parameter will not be a very useful measure of uncertainty. The SE is interpreted as a plus-minus assessment of how sure you are of the parameter value. But if the parameter is very asymmetrical, then a single SE cannot really describe the uncertainty.

• The confidence interval for that parameter cannot be interpreted at face value. Prism always computes the CI from the SE of that parameter (and the number of degrees of freedom), and the CI is always symmetrical around the best-fit value. If the parameter is very asymmetrical, then that symmetrical confidence interval does not give an accurate picture of the uncertainty.

Hougaard’s measure of skewness

Hougaard (1) developed a way to assess the skewness of a parameter used in nonlinear regression without doing any simulations. Prism will
compute this value for each parameter when you check the option the Diagnostics tab of nonlinear regression in the section labeled "Are the parameters intertwined, redundant or skewed?". The results are tabulated along with the other results of nonlinear regression.

Ratkowsky has proposed the following interpretation:

<table>
<thead>
<tr>
<th>Absolute value</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;0.10</td>
<td>Ideal. Almost linear. Confidence intervals can be interpreted at face value</td>
</tr>
<tr>
<td>0.10 - 0.25</td>
<td>Adequate</td>
</tr>
<tr>
<td>0.25 - 1.00</td>
<td>Noticeable skewness. Consider alternative parameterizations of the equation</td>
</tr>
<tr>
<td>&gt; 1.00</td>
<td>Considerable skewness. Strongly consider alternatives</td>
</tr>
</tbody>
</table>

Note that these values are for the absolute value of the Hougaard's measure.

For the simulated data set for the example above, Hougaard's skewness is 0.09 for Khalf and 1.83 for Kprime. This one value (no simulations needed) tells you to choose the form of the model that fits Khalf rather than the form that fits Kprime.

Notes

- Hougaard's measure of skewness is measured for each parameter in the equation (omitting parameters fixed to constant values).

- Prism does not compute Hougaard's skewness if you chose unequal weighting, or a robust fit, because the method is not defined for these situations.

- The values depend on the equation, the number of data points, the spacing of the X values, and the values of the parameters.

- Hougaard's measure of skewness has no units.
• The SAS documentation does a great job of explaining Hougaard's measure (3).

References


3. SAS documentation about Hougaard's measure.

3.5.1.15 Test for appropriate weighting/homoscedasticity

Overview

If you don't change the default weighting, nonlinear regression assumes that, on average, the vertical distance of the points from the curve is the same all the way along the curve. This assumption goes by the name homoscedasticity, and Prism can test this assumption with a test for homoscedasticity.

If you have chosen to differentially weight the points, Prism assumes that the weighted distance of the points from the curve is the same all the way along the curve. Prism tests this assumption with the test for appropriate weighting.

The null hypothesis is that you picked the right weighting scheme, so there is no correlation between the Y value of the curve and the absolute value of the weighted residual. A high P value is consistent with this hypothesis. A small P value suggests that your data violate that assumption. In this case, it might make sense to choose a more appropriate weighting scheme.

Details of the method

To run these tests, Prism follows these steps.
1. For each point, compute the difference between the actual Y value and the Y value of the fit curve at that value of X. This is called a residual.

2. If you chose a weighting scheme, apply this scheme to each residual. If you chose relative weighting, divide each residual by the predicted value of Y. Note a confusing point. In the weighting tab, you choose how to weight the square of the residuals. So the relative weighting is shown in the dialog as dividing by \( Y^2 \). Here we are weighting the residual itself, not the square of the residual, so divide by Y.

3. Compute the absolute values of all the weighted residuals.

4. Create a new table (not shown) where the X values are the absolute values of the weighted residuals and the Y values are the predicted Y values of the curve. There will be one row in this table for each point entered into the regression.

5. Compute the Spearman rank correlation and compute the corresponding P value.

This method has not been published as far as we know, but it is also used by SigmaPlot (where we learned about it). Note that SigmaPlot does exactly what we say above. Even though their manual says they use the observed Y value in step 4, they actually use the predicted value. We don't know of any publication that defines this test. If you search for the Spearman test for hereoscedasticity you'll find that is usually used with linear regression, and where the residuals are correlated with the X value, rather than the predicted Y value.

3.5.1.16 Could the fit be a local minimum?

What is a false minimum?

The nonlinear regression procedure adjusts the variables in small steps in order to improve the goodness-of-fit. If Prism converges on an answer, you can be sure that altering any of the variables a little bit will make the fit worse. But it is theoretically possible that large changes in the variables might lead to a much better goodness-of-fit. Thus the curve that Prism decides is the "best" may really not be the best.
Think of latitude and longitude as representing two variables Prism is trying to fit. Now think of altitude as the sum-of-squares. Nonlinear regression works iteratively to reduce the sum-of-squares. This is like walking downhill to find the bottom of the valley. When nonlinear regression has converged, changing any variable increases the sum-of-squares. When you are at the bottom of the valley, every direction leads uphill. But there may be a much deeper valley over the ridge that you are unaware of. In nonlinear regression, large changes in variables might decrease the sum-of-squares.

This problem (called finding a local minimum) is intrinsic to nonlinear regression, no matter what program you use. You will rarely encounter a local minimum if your data have little scatter, you collected data over an appropriate range of X values, and you have chosen an appropriate equation.

**Testing for a false minimum**

To test for the presence of a false minimum:

1. Note the values of the variables and the sum-of-squares from the first fit.

2. Make a large change to the initial values of one or more parameters and run the fit again.

3. Repeat step 2 several times.

Ideally, Prism will report nearly the same sum-of-squares and same parameters regardless of the initial values. If the values are different, accept the ones with the lowest sum-of-squares.
3.5.1.17 Outliers

**The outliers table**

If you choose to exclude or count outliers as part of nonlinear regression, one of the results tables will list all outliers.

Keep in mind an important distinction:

- If you chose to count outliers (on the Diagnostics tab), then the outliers are still included in the fit.
- If you chose to exclude outliers (on the Fit tab), then the outliers are ignored by the fit, but are still included on the graph.

In both cases, the aggressiveness of the outlier hunt is determined by the ROUT coefficient Q that you enter on the Weights tab.

**What does it mean to be an outlier**

Prism uses the ROUT algorithm, using the value of Q that you can adjust in the Weights tab, to decide when to declare a point to be an outlier. Some considerations:

- The first thing to do is make sure it isn't simply a problem in data entry. If so, fix it.
- Review the list of situations where the outlier identification procedure should not be used.
- Be alert to biological variation. In some experimental situations, outliers can only arise from lab mistakes. These should be excluded. In other situations, an outlier can be the result of biological variation. You may have discovered a new polymorphism in some gene. It could be missing a big opportunity if you exclude and ignore those values.
What to do once you have identified an outlier

After you have identified one or more outliers, you have several choices.

- While outliers are ignored by the nonlinear regression calculation, they are still plotted on the graph. If you want the outlier to be entirely removed from the graph, including error bars on the graph, go back to the data table and exclude the outlier(s). Prism will then refit the curve, and replot the graph without any outliers.

- If you want the outlier to remain on the graph, simply polish the graph Prism provides. Keep in mind that the outliers are a dataset, so you can separately adjust the size, color and symbol of outliers and all the data points. Use the "Datasets on graph" tab of Format Graph to adjust the back-to-front order of data sets to keep the outlier in front of the other data sets. The dialog shows the datasets in back to front order, so you want to keep the outlier data sets lower in the list.

- If you want to remove the special formatting of outliers on the graph, use the "Datasets on graph" tab of Format Graph. Select the outlier dataset(s) and click Remove. The outlier will still be plotted as part of the full data set, but will no longer be plotted separately as an outlier data set.

- If you asked Prism to count the outliers (Diagnostics tab) but now wish to fit the data with the outlier ignored, go back to the nonlinear regression dialog and choose "Automatic outlier elimination" (on the Fit tab).

3.5.1.18 Troubleshooting nonlinear regression

The #1 tip for troubleshooting

Nonlinear regression starts with initial values for each parameter. If these initial values are very far from the correct values, nonlinear regression might go astray. To check this, go to the Diagnostics tab of the nonlinear regression dialog and choose the option at the top of the dialog: Don't fit a curve. Instead plot the curve defined by the initial values of the parameters.

If the curve defined by the initial values does not generally follow the shape of the data, and go near the data points, you should change the initial values (Initial Values tab of nonlinear regression) until they do.
If Prism reports an error code

Prism reports error codes as a single phrase that appears over the column of results, for example "Bad initial values" or "Impossible weights". Learn more about each error message.

Common problems in nonlinear regression

Here is a short list of potential curve fitting problems with suggested solutions.

The equation simply does not describe the data.

Try a different equation.

The initial values are too far from their correct values.

Enter different initial (estimated) values for the parameters. If you entered your own equation, check that you entered sensible rules for generating initial values.

The range of X values is too narrow to define the curve completely.

If possible, collect more data. Otherwise, hold one of the parameters to a constant value.

You have not collected enough data in a critical range of X values.

Collect more data in the important regions.

Your data are very scattered and don't really define a curve.

Try to collect less scattered data. If you are combining several experiments, consider normalizing the data for each experiment to an internal control.

The equation includes more than one component, but your data don't follow a multicomponent model.

Use a simpler equation.
Your numbers are too large or too small.

If your X or Y values are huge, divide by a constant to change the units. Avoid values greater than, say, 100,000.

If your X or Y values are tiny, multiply by a constant change the units. Avoid values less than about 0.00001.

Changing the units is unlikely to solve the problem, but it is worth a try.

You've set a parameter to an inappropriate constant value.

Check that you haven't made a simple mistake like setting a maximum plateau to 1.0 when it should be 100, or a Hill slope to +1.0 when it should be -1.0.

Consult an analysis check list

Prism has three different analysis checklists for nonlinear regression. Use the one that matches your goal.

Analysis checklist: Fitting a model

Analysis checklist: Comparing nonlinear fits

Analysis checklist: Interpolating from a standard curve

3.5.1.19 Why results can differ in various Prism versions

Prism 5 and 6 use the same algorithms, so should always report identical results. Prism 7 has slightly improved algorithms, so the results might vary a bit, but the differences are trivial.

However, Prism 4 used slightly different algorithms, so curve fitting results can differ from results with later Prism versions in these cases:

- If your fit is labeled "Ambiguous" by Prism 5 or later, you know that some of the parameters are not determined precisely. Prism 4 presented a full set of results in this case, but the results are not useful when the fit is ambiguous.
• If you chose no weighting, check the sum-of-squares from the two programs. The goal of regression is to minimize that sum of squares, so see which version of Prism found a fit with the smaller sum-of-squares. Prism 5 and later have a few improvements in the fitting algorithm, so occasionally it can find a better fit than did Prism 4. The differences, if any, are usually trivial.

• If you chose to weight by the Y values (or the Y values squared), Prism 5 and later handle weighting differently than did Prism 4. Prism now weights by the Y value of the curve, while Prism 4 (and earlier releases) weighted by the Y value of the data. Weighting by the Y value of the curve is better, so the results of Prism 5 and later are more correct. Since the weighting is computed differently, you can't directly compare the weighted sum-of-square values reported by the two versions of Prism.

• When you compare two models, Prism now does an extra step. If one of the models is ambiguous, then Prism chooses the other model, without doing the F test or AIC comparison.

• Prism now offers more rules for defining initial parameter values. If your equation uses one of these new rules, Prism 4 might not be able to find a reasonable fit until you tweak those initial values. In particular, Prism now has smarter rules for fitting sigmoidal log(dose) vs. response curves.

• If you entered data as mean, SD (or SEM) and N, then Prism 4 (by default) fits the means and weights by the sample size (N). This is one of the two options on the weighting tab (the other option is to fit means only, ignoring N). Prism is now smarter by default (although you can choose to just fit the means and ignore N and SD values). It accounts not only for sample size N but also for the SD (or SEM) values you enter. With Prism 5 or later (but not Prism 4), you'll get exactly the same results from data entered as mean, SD and N as you would have by entering raw data. Prism 4 only accounts for differences in N, but not SD. The best fit values of the parameters, and thus the appearance of the curve, is the same with Prism 4 and 5. But Prism 5 and later does a smarter job with standard errors, confidence intervals, and comparisons of models.

3.5.2 Interpreting results: Comparing models

In many cases, your primary goal in fitting curves is to compare models, or compare fits of two (or more) data sets.
• First review the general principles of comparing models.

• Next review the method you choose to use: Extra sum-of-squares or AIC model comparison.

• Review how Prism compares models when it detects outliers.

3.5.2.1 Interpreting comparison of models

Reality check before statistics

Apply a common sense reality check before looking at any statistical approach to comparing fits. If one of the fits has results that are scientifically invalid, then accept the other model. Only when both fits make scientific sense should you use statistical method to compare the two fits.

Prism partially automates this 'reality check' approach. If the fit of either model is ambiguous, then Prism chooses the other model without performing any statistical test.

Statistical approaches balance the change in sum-of-squares with the change in numbers of degrees of freedom

The more complicated model (the one with more parameters) almost always fits the data better than the simpler model. Statistical methods are needed to see if this difference is enough to prefer the more complicated model. Prism can do this via the extra sum-of-squares F test or using information theory and computation of AIC. Don't use R² or the adjusted R² to compare models (1).

Both these methods only make sense when the models being compared have different numbers of parameters, and so have different numbers of degrees of freedom. If you want to compare two models with the same number of parameters, there is no need to use either the F test or AIC. Simply choose the model that fits the data the best with the smallest sum-of-squares.
How do these methods work to compare data sets?

The Compare tab of Prism lets you ask "Do the best-fit values of selected unshared parameters differ between data sets?" or "Does one curve adequately fit all data sets?". Applying the F test or Akaike's method to answering these questions is straightforward. Prism compares the sum-of-squares of two fits.

- In one fit, the model is separately fit to each data set, and the goodness-of-fit is quantified with a sum-of-squares. The sum of these sum-of-square values quantifies the goodness of fit of the family of curves fit to all the data sets.

- The other fit is a global fit to all the data sets at once, sharing specified parameters. If you ask Prism whether one curve adequately fits all data sets, then it shares all the parameters.

These two fits are nested (the second is a simpler case of the first, with fewer parameters to fit) so the sums-of-squares (actually the sum of sum of squares for the first fits) can be compared using either the F test or Akaike's method.


3.5.2.2 Interpreting the extra sum-of-squares F test

The extra-sum-of-squares F test compares nested models

The extra-sum-of-squares F test compares the goodness-of-fit of two alternative nested models. "Nested" means that one model is a simpler case of the other. Let's consider what this means in different contexts:

- If you asked Prism to test whether parameters are different between treatments, then the models are nested. You are comparing a model where Prism finds separate best-fit values for some parameters vs. a model where those parameters are shared among data sets. The second case (sharing) is a simpler version (fewer parameters) than the first case (individual parameters).
• If you asked Prism to test whether a parameter value is different than a hypothetical value, then the models are nested. You are comparing the fit of a model where a parameter is fixed to a hypothetical value to the fit of a model where Prism finds the best-fit value of that parameter. The first case (fixed value) is a simpler version (fewer parameters to fit) than the second.

• If you are comparing the fits of two equations you chose, and both models have the same number of parameters, then the two models cannot be nested. With nest models, one model has fewer parameters than the other. When the two models have the same number of parameters, Prism reports that it cannot compute the F test because the two models have the same number of degrees of freedom. In this case, Prism does not report a P value, and chooses to plot the model whose fit has the lower sum-of-squares. Prism reports the error message, "Models have the same DF."

• If you are comparing the fits of two equations you chose with different numbers of parameters, the models may or may not be nested. Prism does not attempt to do the algebra necessary to make this determination. If you chose two models that are not nested, Prism will report results for the extra-sum-of-squares F test, and these results will not be useful.

Interpreting the P value

The extra-sum-of-squares F test is based on traditional statistical hypothesis testing. The F test compares the improvement of SS with the more complicated model vs. the loss of degrees of freedom.

The null hypothesis is that the simpler model (the one with fewer parameters) is correct. The improvement of the more complicated model is quantified as the difference in sum-of-squares. You expect some improvement just by chance, and the amount you expect by chance is determined by the number of degrees of freedom in each model. The F test compares the difference in sum-of-squares with the difference you’d expect by chance. The result is expressed as the F ratio, from which a P value is calculated.

The P value answers this question:

If the null hypothesis is really correct, in what fraction of experiments (the size of yours) will the difference in sum-of-squares be as large as you observed, or even larger?
If the P value is small, conclude that the simple model (the null hypothesis) is wrong, and accept the more complicated model. Usually the threshold P value is set at its traditional value of 0.05.

If the P value is high, conclude that the data do not present a compelling reason to reject the simpler model.

Prism names the null and alternative hypotheses, and reports the P value. You set the threshold P value in the Compare tab of the nonlinear regression dialog. If the P value is less than that threshold, Prism chooses (and plots) the alternative (more complicated) model. It also reports the value of F and the numbers of degrees of freedom, but these will be useful only if you want to compare Prism's results with those of another program or hand calculations.

**When Prism won't report a P value when comparing models**

Prism skips the extra sum of squares test and does not report a P value in these situations:

- If the simpler model fits the data better than (or the same) as the more complicated model. The whole point of the F test is to deal with a tradeoff. The model with more parameters fits the data better fit, but that may just be due to chance. The F test asks if that improvement in fit (decrease in sum of squares) is large enough to be "worth" the loss in degrees of freedom (increase in number of parameters). In the rare cases where the simpler model fits better (or the same) as the more complicated model (the one with more parameters), Prism will choose the simpler model without computing the F test and report "Simpler model fits better".

<table>
<thead>
<tr>
<th>Nonlin fit</th>
<th>A</th>
<th>B</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Table of results</td>
<td>Reference</td>
<td>Sample</td>
<td>Global (shared)</td>
</tr>
<tr>
<td>1</td>
<td>Comparison of Fits</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>2</td>
<td>Null hypothesis</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Alternative hypothesis</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>P value</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>Conclusion (alpha = 0.01)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>Preferred model</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>F (DFr, DFd)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

- If the fit of either model is ambiguous or flagged, then Prism chooses the other model without performing any statistical test. You
have a choice in the Compare tab of nonlinear regression to turn off this criterion.

- If the fit of one model did not converge, then Prism chooses the other model without doing the F test. Since the fit of one model didn't converge, it makes no sense to compare the sum-of-squares of the two models.

- If one model fits the data perfectly, so the sum of squares equals zero. If one model fits perfectly, Prism chooses it without doing the F test.

- If the two models have the same number of degrees of freedom. The idea of the F test is to balance the improvement of sum-of-squares (better fit) with the decrease in degrees of freedom (more parameters). The F test makes no sense (and is mathematically impossible due to division by zero) if the two models have the same number of degrees of freedom. In this case, Prism picks the model that fits best.

3.5.2.3 Interpreting AIC model comparison

This alternative approach is based on information theory, and does not use the traditional “hypothesis testing” statistical paradigm. Therefore it does not generate a P value, does not reach conclusions about “statistical significance”, and does not “reject” any model.

The method determines how well the data supports each model, taking into account both the goodness-of-fit (sum-of-squares) and the number of parameters in the model. The results are expressed as the probability that each model is correct, with the probabilities summing to 100%. If one model is much more likely to be correct than the other (say, 1% vs. 99%), you will want to choose it. If the difference in likelihood is not very big (say, 40% vs. 60%), you will know that either model might be correct, so will want to collect more data.

Of course, these probabilities are meaningful only in the context of comparing those two models. It is possible a third model you didn't test fits much better so is much more likely to be correct.
Prism names the null and alternative hypotheses, and reports the likelihood that each is correct. It also reports the difference between the AICc values (as the AICc of the simple model minus the AICc of the more complicated model), but this will be useful only if you want to compare Prism's results with those of another program or hand calculations. Prism chooses and plots the model that is more likely to be correct, even if the difference in likelihoods is small.

### 3.5.2.4 How Prism compares models when outliers are eliminated

Comparing models only makes sense when both models 'see' exactly the same set of data. That makes it tricky to combine outlier elimination with model comparison. Prism handles this situation automatically.

If you select automatic outlier elimination (on the Fit tab), Prism first fits each model separately to determine which points (if any) are outliers. Then it removes all the points that are defined as outliers from either of the models, and fits both models again. It uses the sum-of-squares and df from these fits to perform the comparison. This approach ensures that exactly the same points are fit to both models, and that outliers from either model are excluded.

### 3.5.2.5 Interpreting the adjusted $R^2$

**Why you should not use $R^2$ to compare models**

$R^2$ quantifies how well a model fits the data, so it seems as though it would be an easy way to compare models. It sure sounds easy -- pick the model with the larger $R^2$. The problem with this approach is that there is no penalty for adding more parameters. So the model with more parameters will bend and twist more to come nearer the points, and so almost always has a higher $R^2$. If you use $R^2$ as the criteria for picking the best model, you'd almost always pick the model with the most parameters.

**The adjusted $R^2$ accounts for the number of parameters fit**

The adjusted $R^2$ always has a lower value than $R2$ (unless you are fitting only one parameter). The equations below show why.
The equations above show how the adjusted $R^2$ is computed. The sum-of-squares of the residuals from the regression line or curve have $n-K$ degrees of freedom, where $n$ is the number of data points and $K$ is the number of parameters fit by the regression. The total sum-of-squares is the sum of the squares of the distances from a horizontal line through the mean of all Y values. Since it only has one parameter (the mean), the degrees of freedom equals $n-1$.

When $K=1$, adjusted $R^2$ and the ordinary $R^2$ are identical. When $K>1$, The adjusted $R^2$ is smaller than the ordinary $R^2$ (the Prism 6 Help had this backwards).

Using adjusted $R^2$ and a quick and dirty way to compare models

A quick and easy way to compare models would seem to be to choose the one with the smaller adjusted $R^2$. Choose to report this value on the Diagnostics tab.

Comparing models with adjusted $R^2$ is a standard method for comparing models fit with multiple linear regression. It is not standard for nonlinear regression for good reason. Speis and Neumeyer(1) did simulations that showed that using the adjusted $R^2$ is a horrible way to choose among competing models. We suggest that you use the extra-sum-of-square F test or comparing AICc instead. If you do compare models by comparing adjusted $R^2$, make sure that identical data, weighted identically, are used for all fits. But even then, know that models that fit very differently as assessed by AICc may have $R^2$ values that differ only in the third to fifth digit after the decimal (1).
Adjusted $R^2$ in linear regression

Prism doesn't report the adjusted $R^2$ with linear regression, but you can fit a straight line with nonlinear regression.

If X and Y are not linearly related at all, the best fit slope is expected to be 0.0. If you analyzed many randomly selected samples, half the samples would have a slope that is positive and half the samples would have a negative slope. But in all these cases, $R^2$ would be positive (or zero). $R^2$ can never be negative (unless you constrain the slope or intercept so it is forced to fit worse than a horizontal line). In contrast, the adjusted $R^2$ can be negative. If you analyzed many randomly selected samples, you'd expect the adjusted $R^2$ to be positive in half the samples and negative in the other half.

Here is a simple way to think about the distinction. The $R^2$ quantifies the linear relationship in the sample of data you are analyzing. Even if there is no underlying relationship, there almost certainly is some relationship in that sample. The adjusted $R^2$ is smaller than $R^2$ and is your best estimate of the degree of relationship in the underlying population.


3.5.2.6 Comparing fits with ANOVA

What if you wish to test for differences in best-fit parameters between three or more data sets, on the basis of a single experiment? You don't just want to know if all the curves are the same. You want to use multiple comparisons to compare pairs of curves, focusing on a particular parameter. Prism lets you do this in two ways:

ANOVA approach -- statistical significance

1. Perform the nonlinear regression analysis. Record from the Results sheet the best-fit values for the parameter you are comparing, perhaps the logEC50 of a dose response curve.

2. Also record the standard errors for those parameters and the degrees of freedom for each curve (which equals the number of data points minus the number of variables fit).
3. Create a new Grouped table in Prism, formatted for entry of "Mean, Standard Error, N". You will enter values only into the first row of this table.

4. For each data set, enter the best-fit value of the parameter (i.e. logEC50) in the "Mean" column.

5. Enter the standard error of the best-fit value in the "SEM" column.

6. For N, enter one more than the degrees of freedom for that fit. (Why enter df+1 into the "N" column? The ANOVA calculations don't actually care about the value of N. Instead, they are based on df. Prism subtracts 1 from the value you enter as N, and uses that as dfF. Since you enter df+1, Prism ends up using the correct df value.)

7. Click Analyze and choose one-way ANOVA along with an appropriate multiple comparisons test.

**Compare two curves at a time with nonlinear regression**

You can rerun the analysis comparing two data sets (curves) at a time. The easiest way to do this is to duplicate the results of the main analysis (New..Duplicate sheet) and then remove all but two data sets from that new analysis. Another approach is to keep one data table, click Analyze, choose nonlinear regression and on the right side of that dialog choose which two data sets to compare.

There are two approaches to use when comparing fits, the extra sum-of-squares F test and the AICc approach.

With statistical significance (extra sum-of-squares F test) approach, there is a traditional (albeit totally arbitrary) cutoff at P=0.05. But if you are doing many comparisons, you should correct for the multiple comparisons. Divide 0.05 (or whatever overall value you want) by the number of pairs of analyses you are comparing, to come up with a new stricter cut off for declaring a P value to be small enough that you can call the comparison "significant".

The AIC approach to comparing curves is not based on statistical hypothesis testing, and is not confused by multiple comparisons. There are two ways to use this approach:
In the Diagnostics tab of nonlinear regression of Prism 6, check the option to report the AICc of each curve. Then you can do manual calculations or comparisons with those AICc values.

Run the nonlinear regression with two data sets at a time, and use the AIC approach to ask how strong the evidence is that the parameter you care about (logEC50) differs between data sets. The AIC calculations just give you the likelihood that the parameter is the same in both data sets vs. different. You need to decide when those likelihood are far enough apart that you will believe the parameters are different.

3.5.3 Analysis checklists: Nonlinear regression

To understand your results, review the appropriate analysis checklist, depending on your experimental goal.

- **Analysis checklist: Fitting a model**
- **Analysis checklist: Comparing nonlinear fits**
- **Analysis checklist: Interpolating from a standard curve**

3.5.3.1 Analysis checklist: Fitting a model

Your approach in evaluating nonlinear regression depends on your goal.

In many cases, your goal is to create a standard curve from which to interpolate unknown values. We've created a different checklist for this purpose.

More often, your goal is to determine the best-fit values of the model. If that is your goal, here are some questions to ask yourself as you evaluate the fit:
Curve

✅ Does the graph look sensible?

Your first step should be to inspect a graph of the data with superimposed curve. Most problems can be spotted that way.

✅ Does the runs or replicate test tell you that the curve deviates systematically from the data?

The runs and replicates tests are used to determine whether the curve follows the trend of your data. The runs test is used when you have single Y values at each X. It asks if data points are clustered on either side of the curve rather than being randomly scattered above and below the curve. The replicates test is used when you have replicate Y values at each X. It asks if the points are 'too far' from the curve compared to the scatter among replicates.

If either the runs test or the replicates test yields a low P value, you can conclude that the curve doesn't really describe the data very well. You may have picked the wrong model, or applied invalid constraints.

Parameters

✅ Are the best-fit parameter values plausible?

When evaluating the parameter values reported by nonlinear regression, check that the results are scientifically plausible. Prism doesn't 'know' what the parameters mean, so can report best-fit values of the parameters that make no scientific sense. For example, make sure that parameters don't have impossible values (rate constants simply cannot be negative). Check that EC50 values are within the range of your data. Check that maximum plateaus aren't too much higher than your highest data point.

If the best-fit values are not scientifically sensible, then the results won't be useful. Consider constraining the parameters to a sensible range, and trying again.

✅ How precise are the best-fit parameter values?
You don't just want to know what the best-fit value is for each parameter. You also want to know how certain that value is. Therefore an essential part of evaluating results from nonlinear regression is to inspect the 95% confidence intervals for each parameter.

If all the assumptions of nonlinear regression are true, there is a 95% chance that the interval contains the true value of the parameter. If the confidence interval is reasonably narrow, you've accomplished what you wanted to do – found the best fit value of the parameter with reasonable certainty. If the confidence interval is really wide, then you've got a problem. The parameter could have a wide range of values. You haven't nailed it down. How wide is 'too wide' depends on the scientific context of your work.

Are the confidence bands 'too wide'?

Confidence bands visually show you how precisely the parameters have been determined. Choose to plot confidence bands by checking an option on the Fit tab of the nonlinear regression dialog. If all the assumptions of nonlinear regression have been met, then there is a 95% chance that the true curve falls between these bands. This gives you a visual sense of how well your data define the model.

Residuals

Does the residual plot look good?

A residual plot shows the relationship between the X values of your data and the distance of the point from the curve (the residuals). If the assumptions of the regression are met, the residual plot should look bland, with no trends apparent.

Does the scatter of points around the best-fit curve follow a Gaussian distribution?

Least squares regression is based on the assumption that the scatter of points around the curve follows a Gaussian distribution. Prism offers three normality tests (in the Diagnostics tab) that can test this assumption (we recommend the D'Agostino test). If the P value for a normality test is low, you conclude that the scatter is not Gaussian.

Could outliers be impacting your results?
The presence of one or a few outliers (points much further from the curve than the rest) can overwhelm the least-squares calculations and lead to misleading results.

You can spot outliers by examining a graph (so long as you plot individual replicates, and not mean and error bar). But outliers can also be detected automatically. GraphPad has developed a new method for identifying outliers we call the ROUT method. Check the option on the diagnostics tab to count the outliers, but leave them in the calculations. Or check the option on the Fit tab to exclude outliers from the calculations.

Models

✓ Would another model be more appropriate?

Nonlinear regression finds parameters that make a model fit the data as closely as possible (given some assumptions). It does not automatically ask whether another model might work better.

Even though a model fits your data well, it may not be the best, or most correct, model. You should always be alert to the possibility that a different model might work better. In some cases, you can't distinguish between models without collecting data over a wider range of X. In other cases, you would need to collect data under different experimental conditions. This is how science moves forward. You consider alternative explanations (models) for your data, and then design experiments to distinguish between them.

✓ If you chose to share parameters among data sets, are those datasets expressed in the same units?

Global nonlinear regression (any fit where one or more parameter is shared among data sets) minimizes the sum (over all datasets) of the sum (over all data points) of the squared distance between data point and curve. This only makes sense if the Y values for all the datasets are expressed in the same units.

Goodness of fit

✓ Is the $R^2$ 'too low' compared to prior runs of this experiment?
While many people look at $R^2$ first, it really doesn't help you understand the results very well. It only helps if you are repeating an experiment you have run many times before. If so, then you know what value of $R^2$ to expect. If the $R^2$ is much lower than expected, something went wrong. One possibility is the presence of outliers.

Are the values of sum-of-squares and sy.x 'too low' compared to prior runs of this experiment?

These values are related to the $R^2$, and inspecting the results can only be useful when you have done similar experiments in the past so know what values to expect.

3.5.3.2 Analysis checklist: Comparing nonlinear fits

Approach to comparing models

Which model is 'best'? At first, the answer seems simple. The goal of nonlinear regression is to minimize the sum-of-squares, so the model with the smaller sum-of-squares clearly fits best. But that is too simple.

The model with fewer parameters almost always fits the data worse. When you compare two models, the simpler model (with fewer parameters) will have a higher sum-of-squares. When you compare datasets, the global fit will have a higher sum-of-squares than the sum of the individual fits. And when you compare fits with and without a parameter constrained to a particular value, will have a higher sum-of-squares than a model where you don't constrain the parameter.

Both the F test and the AIC method take into account both the difference in goodness-of-fit but also the number of parameters. The question is whether the decrease in sum of squares (moving from simpler to more complicated model) is worth the 'cost' of having more parameters to fit. The F test and AICc methods approach this question differently.

How Prism reports model comparison

Prism reports the results for the comparison at the top of the nonlinear regression table of results. It clearly states the null and alternative hypotheses. If you chose the F test, it reports the F ratio and P value. If
If you chose Akaike's method, it reports the difference in AICc and the probability that each model is correct.

<table>
<thead>
<tr>
<th></th>
<th>Comparison of Fits</th>
<th></th>
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<tbody>
<tr>
<td>1</td>
<td>Comparison of Fits</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Null hypothesis</td>
<td>Polynomial: First Order (straight line)</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Alternative hypothesis</td>
<td>Polynomial: Second Order ($Y = A + B \times X + C \times X^2$)</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>F value</td>
<td>0.0005</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>Conclusion (alpha = 0.05)</td>
<td>Reject null hypothesis</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>Preferred model</td>
<td>Polynomial: Second Order ($Y = A + B \times X + C \times X^2$)</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>F (DFn, DFd)</td>
<td>37.32 (1,7)</td>
<td></td>
</tr>
</tbody>
</table>

Finally, Prism reports the 'preferred' model. You should understand how Prism decides which model is 'preferred' as you may 'prefer' the other model.

If you chose the extra sum-of-squares F test, then Prism computes a P value that answers this question:

If the null hypothesis is really correct, in what fraction of experiments (the size of yours) will the difference in sum-of-squares be as large as you observed or larger?

In the Compare tab, you also tell Prism which P value to use as the cutoff (the default is 0.05). If the P value is lower than that value, Prism chooses the more complicated model. Otherwise it chooses the simpler model.

If you chose Akaike's method, Prism chooses the model that is more likely to be correct. But you should look at the two probabilities. If they are similar in value, then the evidence is not persuasive and both models fit pretty well.

**Analysis checklist for comparing models**

**Was it possible to fit both models?**

Before running the extra sum-of-squares F test or computing AICc values, Prism first does some common sense comparisons. If Prism is unable to fit either model, or if either fit is ambiguous[261] or perfect[264], then Prism does not compare models.
✓ Are both fits sensible?

Apply a reality check before accepting Prism's results. If one of the fits has results that are scientifically invalid, then accept the other model regardless of the results of the F test or AIC comparison.

For example, you would not want to accept a two-phase exponential model if the magnitude of one of the phases is only a tiny fraction of the total response (especially if you don’t have many data points). And you wouldn't want to accept a model if the half-life of one of the phases has a value much less than your first time point or far beyond your last time point.

Analysis checklist for the preferred model

Before accepting the 'preferred' model, you should think about the same set of questions as you would when only fitting one model. You can consult the detailed checklist or the abbreviated version below.

✓ How precise are the best-fit parameter values?
✓ Are the confidence bands 'too wide'?
✓ Does the residual plot look good?
✓ Does the scatter of points around the best-fit curve follow a Gaussian distribution?

3.5.3.3 Analysis checklist: Interpolating from a standard curve

Your approach in evaluating nonlinear regression depends on your goal.

In many cases, your goal is to learn from the best-fit values. If that is your goal, view a different checklist.
If your goal is to create a standard curve from which to interpolate unknown values, your approach depends on whether this is a new or established assay.

**Established assay**

If the assay is well established, then you know you are fitting the right model and know what kind of results to expect. In this case, evaluating a fit is pretty easy.

- **Does the curve go near the points?**
- **Is the R² 'too low' compared to prior runs of this assay?**

If so, look for outliers, or use Prism's automatic outlier detection.

- **Are the confidence bands too wide?**

The confidence bands let you see how accurate interpolations will be, so we suggest always plotting prediction bands when your goal is to interpolate from the curve. If you are running an established assay, you know how wide you expect the prediction bands to be.

**New assay**

With a new assay, you also have to wonder about whether you picked an appropriate model.

- **Does the curve go near the points?**

Look at the graph. Does it look like the curve goes near the points.

- **Are the prediction bands too wide?**

How wide is too wide? The prediction bands show you how precise interpolations will be. Draw a horizontal line somewhere along the curve, and look at the two places where that line intercepts the prediction bands. This will be the confidence interval for the interpolation.

- **Does the scatter of points around the best-fit curve follow a Gaussian distribution?**
Least squares regression is based on the assumption that the scatter of points around the curve follows a Gaussian distribution. Prism offers three normality tests (in the Diagnostics tab) that can test this assumption (we recommend the D'Agostino test). If the P value for a normality test is low, you conclude that the scatter is not Gaussian.

✓ **Could outliers be impacting your results?**

Nonlinear regression is based on the assumption that the scatter of data around the ideal curve follows a Gaussian distribution. This assumption leads to the goal of minimizing the sum of squares of distances of the curve from the points. The presence of one or a few outliers (points much further from the curve than the rest) can overwhelm the least-squares calculations and lead to misleading results.

You can spot outliers by examining a graph (so long as you plot individual replicates, and not mean and error bar). But outliers can also be detected automatically. GraphPad has developed a new method for identifying outliers we call the ROUT method. Check the option on the diagnostics tab to count the outliers, but leave them in the calculations. Or check the option on the Fit tab to exclude outliers from the calculations.

✓ **Does the curve deviate systematically from the data?**

If either the runs test or the replicates test yields a low P value, then you can conclude that the curve doesn't really describe the data very well. You may have picked the wrong model.

### 3.5.4 Error messages from nonlinear regression

When Prism hits a problem, it will present a very brief message in the top row of the results tables. Read the pages in this section to learn what these messages mean.

"Bad initial values" [257]

"Interrupted" [257]
"Not converged"

"Ambiguous"

"Hit constraint"

"Don't fit"

"Too few points"

"Perfect fit"

"Impossible weights"

"Equation not defined"

3.5.4.1 "Bad initial values"

What does "bad initial values" mean?

Nonlinear regression works iteratively. Prism starts with initial estimated values for each parameter. It then gradually adjusts these until it converges on the best fit. In rare circumstances, Prism cannot compute a curve based on the initial values you entered. When it tries, it encounters a math problem such as division by zero or taking the logarithm of a negative number. In this case, Prism does not report any results and instead reports "bad initial values".

Checklist

✓ Did you enter or choose the right equation?

✓ Did you enter sensible values for initial values?

✓ Is the range of X values invalid? Focus on the first and last X value (and on X=0 if included in the range). Can the equation be evaluated at those X values with the initial values you entered?
3.5.4.2 "Interrupted"

What does 'interrupted' mean?

Prism reports 'interrupted' in two situations:

- Nonlinear regression is iterative and the maximum number of iterations is specified in the Diagnostics tab, with 1000 being the default. With data that doesn't really define all the parameters in the model, Prism can just keep iterating until it hits the limit. In that case it stops and reports "interrupted". You can lift the limit in the Diagnostics tab of the nonlinear regression dialog, but usually the problem is that the data simply doesn't enough information for the model to fit.

- The fit was slow, and you clicked "Interrupt" on the progress dialog.

Checklist

If the maximum number of iterations was set to a low value, set it to a higher value and try again. If you have lots of data points and lots of parameters, nonlinear regression can sometimes require hundreds of iterations.

If the maximum number of iterations was already set to a high value, you can try a still higher value, but most likely Prism is still not going to be able to find a best-fit curve. Things to check:

✔ Did you enter the right model?

✔ Does your data provide enough information to define that model. For example if you fit a log(dose) response curve, does your data show a sigmoidal shape, enough to define the top or bottom plateaus? If your data doesn't define all the parameters, consider constraining one or more parameters to constant values.

✔ Does the curve defined by your initial values come near your data? Check the option on the diagnostics tab to plot that curve.

✔ If you entered constraints, were they entered correctly?
If you didn't enter any constraints, consider whether you can constrain one or more parameters to a constant value? For example, in a dose-response curve can you constrain the bottom plateau to be zero?

Can you share a parameter over all the data sets (global fitting)?

3.5.4.3 "Not converged"

What does "not converged" mean?

Nonlinear regression works iteratively. Prism starts with initial estimated values for each parameter. It then gradually adjusts these until it converges on the best fit. "Converged" means that any small change in parameter values creates a curve that fits worse (higher sum-of-squares). But in some cases, it simply can't converge on a best fit, and gives up with the message 'not converged'. This happens in two situations:

- The model simply doesn't fit the data very well. Perhaps you picked the wrong model, or applied the wrong constraints.

- The initial values generated a curve that didn't come close to the points. In that case, Prism may not be able to figure out how to change the parameters to make the curve fit well.

Checklist

✅ Did you enter the right model?

✅ Does the curve defined by your initial values come near your data? Check the option on the [diagnostics tab to plot that curve.

✅ If you entered constraints, were they entered correctly?

✅ If you didn't enter any constraints, consider whether you can constrain one or more parameters to a constant value? For example, in a dose-response curve can you constrain the bottom plateau to be zero?

✅ Can you share a parameter over all the data sets (global fitting)?
3.5.4.4 ambiguous

3.5.4.5 "Ambiguous"

What does "ambiguous" mean?

"Ambiguous" is a term coined by GraphPad to describe a fit that doesn't really nail down the values of all the parameters.

Changing the value of any parameter will always move the curve further from the data and increase the sum-of-squares. But when the fit is 'ambiguous', changing other parameters can move the curve so it is near the data again. In other words, many combinations of parameter values lead to curves that fit equally well.

Prism puts the word 'ambiguous' in the top row of results. For the parameters that are 'ambiguous' (there may be one or several), Prism puts '~' before the best fit values and standard errors, and reports "very wide" for the corresponding confidence intervals.

If you check the option in the diagnostics tab to report dependency, the value will be >0.9999 for all the 'ambiguous' parameters (that is how we define 'ambiguous'; the threshold value is arbitrary).

If the fit is 'ambiguous' you really can't interpret the best-fit values of some parameters.

Does it matter?

If your goal is to interpolate unknowns from a standard curve, you won't care that the parameter values are 'ambiguous'. So long as the curve goes through the points, and doesn't wiggle too much, the interpolations will be useful.

If your goal is to learn about your data by inspecting the values of the parameters, then you've got a real problem. At least one of the parameters has a best-fit value that you should not rely upon.

But the $R^2$ is really high...

In many cases, the $R^2$ will be really high, maybe 0.99. That just means that the curve comes close to the data points. It doesn't mean the data
define all the parameters. If the fit is 'ambiguous', you can get an equally well-fitting curve with a different set of values of the parameters.

**Reasons for ambiguous fits**

**Data not collected over a wide enough range of X values**

The data above show a fit of a dose-response curve to a set of data that don't define a top plateau. Since the top plateau was not constrained to a constant value, Prism reports the fit to be ambiguous.

**Model too complicated for the data**
The data above fit fine to a standard dose-response curve. But if you try to fit it to a biphasic dose-response curve, Prism reports that the results are ambiguous. The data follow a standard dose-response model just fine, with no evidence of a second component. So fitting to a biphasic model -- with two EC50 values -- is ambiguous.

Model has redundant parameters

The simplest example would be fitting this model: \( Y = A + B + C \times X \). This model describes a straight line with a slope equal to \( C \) and a Y intercept equal to the sum of \( A \) and \( B \). But there is nothing in the data to fit \( A \) and \( B \) individually. There are an infinite number of pairs of values of \( A \) and \( B \) that lead to the same sum, so the same Y intercept. If Prism attempts to fit this model, it will conclude that the fit is ambiguous.

Prism cannot tell the difference between this case and the previous one. But the two cases are distinct. The model in this example has redundant parameters. It doesn't matter what the data look like, or how many data points you collect, fitting this model will always result in 'ambiguous' results, because two of the parameters are intertwined. In the previous example, the fit was ambiguous with the example data set, but would not have been ambiguous with other data sets.

Checklist

- ✔ Can you constrain any parameters to a constant value?
- ✔ Can you share one or more parameters among datasets?
- ✔ Can you collect more data points, perhaps over a wider range of \( X \) values?
- ✔ If your model has two phases or components, consider switching to a one phase or component model.

3.5.4.6 "Hit constraint"

What does "hit constraint" mean?

Prism reports "Hit constraint" when the best-fit value of one or more parameters is right at the limit of a constraint.
When a fit hits a constraint, the results are unlikely to provide useful information. If you had a solid reason to constrain a parameter within a range of values, it ought to end up in that range. If the fit hit the constraint limit, that means the true best-fit value is some value forbidden by the constraint.

Prism does not compute confidence and prediction bands when a parameter hit a constraint. The best-fit values are not a local minimum, so any attempt to compute confidence or prediction bands would give misleading results. Prism does compute the confidence intervals for the other parameters (the ones that didn't hit a constraint) but these need to be viewed with caution.

When a fit ends up hitting a constraint, it is likely that you set the constraint incorrectly. So the first thing to do is make sure the constraint is sensible and correctly entered. For example, if you mistakenly set a constraint that a rate constant K must be less than or equal to zero, Prism's nonlinear regression might converge with the best-fit value equalling zero and report that you hit the constraint. Fix the constrain to be greater than or equal to zero, and the fit will work fine.

Another possibility is to change the constraint from an inequality (Bottom>0) to a constant constraint (Bottom=0). You'll get the same parameter values, but difference confidence intervals, and you can get confidence and prediction bands.

**Checklist**

- Did you enter the constraint correctly? Did you mix up "<" and ">".
- Did you enter the model correctly?
- Are you asking Prism to do the impossible -- too many parameters for your data?
- If your model has two phases or components, consider switching to a one phase or component model.
3.5.4.7 "Don't fit"

The top of the Diagnostics tab offers the choice to plot the curve defined by the initial values of the parameters.

When you make this choice, the top of the results will show "Don't fit".

This is a very useful option to use when diagnosing problem fits. But you won't learn anything by looking at the results page. Instead, look at the graph. If the curve goes near the points, then the initial values are OK. If the curve is far from the points, or follows a different shape altogether, then you know that there is a problem with the choice of model or initial values.

If you checked this option by mistake, go back to the Diagnostics tab and check the alternative choice ("Fit the curve.").

3.5.4.8 "Too few points"

Regression requires at least as many data points as parameters. So if you fit a dose-response model with four parameters (Bottom, Top, logEC50, and HillSlope), you must have four or more points or Prism will simply report "too few points".

Having as many points as parameters is the bare minimum. Prism will try to fit the model, and will report a perfect fit, and so not be able to report standard errors or confidence intervals. If there are only a few more points that parameters, Prism is likely to end up with an ambiguous or not converged fit.

3.5.4.9 "Perfect fit"

Prism reports "perfect fit' when the curve goes through every point. The sum-of-squares is 0.0, and R² is 1.00.

If you are testing nonlinear regression with made up values, add some random scatter to make a better example.
If you are fitting actual data, and the fit is perfect, you either got very lucky or have very few data points. It is not possible to compute the standard errors and confidence intervals of the parameters when the fit is perfect, nor is it possible to compare models.

3.5.4.10 "Impossible weights"

Prism reports "Impossible weighting" when weighting would lead to division by zero for one or more point. This can happen when you choose to weight by the reciprocal of Y, and the Y value of the curve equals zero at some X values. Since division by zero is impossible, Prism reports "Impossible weighting". Go to the weights tab and choose a different weighing method.

Prism also reports "impossible weights" when the Y value of the curve is negative for some values of X. If you weight by 1/Y, this would mean the weights would be negative. If you weight by 1/Y^2, the weights would all be positive, but points with Y near zero would get lots of weight, and points with Y values far from zero (in either direction) would get low weights, and this doesn't really make sense.

3.5.4.11 "Equation not defined"

When you enter a user-defined equation, you can write a different model for different data sets. If no model is defined for certain data sets, the message "Equation not defined" will appear in the results for that data set, with no other results.

- "Bad initial values"
- "Interrupted"
- "Not converged"
- "Ambiguous"
- "Hit constraint"
- "Don't fit"
- "Too few points"
- "Perfect fit"
3.5.4.12 "Can't calculate"

If you ask Prism to use either the F test or the AICc method to compare two models, it will report "Can't calculate" if the two models have the same number of parameters. This means it can't calculate the comparison between fits, not that it can't fit the two models to the data.

Both these methods to compare models only make sense when the models being compared have different numbers of parameters, and so have different numbers of degrees of freedom. These methods are alternative methods of dealing with the tradeoff between how well the model fits and how complicated it is (assessed by the number of parameters).

If you want to compare two models with the same number of parameters, there is no need to use either the F test or AIC. Simply choose the model that fits the data the best with the smallest sum-of-squares.

3.6 Models (equations) built-in to Prism

To use nonlinear regression, you need to choose a model. In many cases, especially if you are fitting pharmacological or physiological data, you can choose one of Prism's built-in models described here.

- **Dose-response - Key concepts**
- **Dose-response - Stimulation**
- **Dose-response - Inhibition**
- **Dose-response -- Special**
- **Receptor binding - Key concepts**
- **Receptor binding - Saturation binding**
- **Receptor binding - Competitive binding**
3.6.1 Dose-response - Key concepts

3.6.1.1 What are dose-response curves?

What are dose-response curves?

Dose-response curves can be used to plot the results of many kinds of experiments. The X axis plots concentration (or dose) of a drug or hormone. The Y axis plots response, which could be almost any measure of biological function.

The term “dose” is often used loosely. In its strictest sense, the term only applies to experiments performed with animals or people, where you administer various doses of drug. You don't know the actual concentration of drug at its site of action—you only know the total dose that you administered. However, the term “dose-response curve” is also used more loosely to describe in vitro experiments where you apply known concentrations of drugs. The term “concentration-response curve” is a more precise label for the results of these types of experiments. The term “dose-response curve” is occasionally used even more loosely to
refer to experiments where you vary levels of some other variable, such as temperature or voltage.

**X values are logarithm of doses or concentrations**

Dose-response experiments typically use around 5-10 doses of agonist, equally spaced on a logarithmic scale. For example, doses might be 1, 3, 10, 30, 100, 300, 1000, 3000, and 10000 nM. When converted to logarithms (and rounded a bit), these values are equally spaced: 0.0, 0.5, 1.0, 1.5, 2.0, 2.5, 3.0, 3.5, and 4.0.

All the dose-response curves built in to Prism are provided in two forms. One form assumes that X is the logarithm of concentration or the logarithm of dose. The other form assumes X is concentration or dose. Make sure you pick the form of the equation to match the data you are analyzing.

The uncertainty in the value of the EC50 or IC50 is usually quite asymmetrical. So be sure to choose the profile likelihood method for computing confidence intervals that produces asymmetrical intervals.

**Y values are responses**

In a dose-response curve, the Y values are responses. For example, the response might be enzyme activity, accumulation of an intracellular second messenger, membrane potential, secretion of a hormone, change in heart rate, or contraction of a muscle.

You can transform the Y values to new units by multiplying or dividing by a constant. Use Prism's Transform analysis for this. Transforming to new units will not fundamentally change the results of a curve fit.

In some cases, the transform from experimentally observed units to practical units is nonlinear. For example, a nonlinear transform is needed to convert the ratio of two fluorescence values to concentrations of Ca++. Which Y values should be used when fitting a dose-response curve? Nonlinear regression assumes that all scatter around the curve is Gaussian, so you want to use whatever units make that assumption most true. In many cases, this may be hard to know.

3.6.1.2 **The terms "agonist", "antagonist", and "normalized" in equation names**

While the terms agonist and antagonist have rigorous meanings in
pharmacology, we use the terms more loosely when naming built-in dose-response equations.

We use the term *agonist* to describe curves that go uphill, with a larger Y value as the X value gets larger.

We use the term *antagonist* to describe curves that go downhill, with a smaller Y value as the X value gets larger.

We use the term *normalized* to describe curves that go between Y=0.0 and Y=100.0. Even if you have normalized your data, you may not want to use the normalized equation. One reason not to use the normalized equation is to let Prism fit the top and/or bottom plateaus. Another reason is that with some drugs (a partial agonist) the top plateau won't be 100 (assuming you define 100 based on the response to a different drug).

3.6.1.3 Converting concentration to log(concentration)

Prism provides two sets of dose-response equations. One set assumes you have entered concentrations (or doses) as X. The other set assumes you entered the logarithms of concentration or dose.

If you entered actual doses or concentrations, instead of their logarithms, but want to use an equation that assumes the X values are logarithms, use Prism to transform the X values.

1. From the data table, click Analyze and then choose Transform Concentrations (X), which is the second analysis listed on the Analyze dialog.

2. On the dialog, check the option to transform X to logarithms. Base 10 (common) logarithms are used most commonly.

3. If any of your X values are 0.0, use the option at the top of the dialog to change these to some small concentration. The reason for this is that the logarithm of zero is not defined.
3.6.1.4 The EC50

What is the EC50?

The EC50 is defined quite simply as the concentration of agonist that provokes a response halfway between the baseline (Bottom) and maximum response (Top). It is impossible to define the EC50 until you first define the baseline and maximum response.

Depending on how you have normalized your data, the EC50 may not be the same as the concentration that provokes a response of Y=50. For instance, in the example below, the data are normalized to percentage of maximum response, without subtracting a baseline. The baseline is about 20%, and the maximum is 100%, so the EC50 is the concentration of agonist that evokes a response of about 60% (halfway between 20% and 100%). The concentration that provokes a response of 50 in this experiment is not the EC50.
Don't over interpret the EC50. It is simply the concentration of agonist required to provoke a response halfway between the baseline and maximum responses. Because the EC50 defines the location of the dose-response curve for a particular drug, it is the most commonly used measure of an agonist’s potency. However, the EC50 is usually not the same as the Kd for the binding of agonist to its receptor -- it is not a direct measure of drug affinity.

The pEC50

The pEC50 is defined as the negative logarithm of the EC50. If the EC50 equals 1 micromolar ($10^{-6}$ molar), the log EC50 is $-6$ and the pEC50 is 6. There is no particular advantage to expressing potency this way, but it is customary in some fields.

The IC50

In many experiments, you vary the concentration of an inhibitor. With more inhibitor, the response decreases, so the dose-response curve goes downhill. With such experiments, the midpoint is often called the IC50 ("I" for inhibition) rather than the EC50 ("E" for effective). This is purely a difference in which abbreviation is used, with no fundamental difference.

ECanything

A simple rearrangement of the equation lets you fit EC80 (or EC90 or ECanything) instead of the EC50. Prism includes an equation that fits any EC value directly.
3.6.1.5 Confidence intervals of the EC50

The 95% confidence interval of the EC50 is not symmetrical

Sample data

<table>
<thead>
<tr>
<th>[Agonist], M</th>
<th>Response</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.00e-009</td>
<td>29</td>
</tr>
<tr>
<td>1.00e-008</td>
<td>68</td>
</tr>
<tr>
<td>3.00e-008</td>
<td>32</td>
</tr>
<tr>
<td>1.00e-007</td>
<td>52</td>
</tr>
<tr>
<td>3.00e-007</td>
<td>154</td>
</tr>
<tr>
<td>1.00e-006</td>
<td>171</td>
</tr>
<tr>
<td>3.00e-006</td>
<td>201</td>
</tr>
<tr>
<td>1.00e-005</td>
<td>341</td>
</tr>
<tr>
<td>3.00e-005</td>
<td>341</td>
</tr>
<tr>
<td>1.00e-004</td>
<td>401</td>
</tr>
<tr>
<td>3.00e-004</td>
<td>389</td>
</tr>
</tbody>
</table>

Asymmetrical confidence intervals
The sample data above were fit to the model: [Agonist] vs. response -- variable slope with the Confidence tab set to compute confidence intervals using the asymmetrical (likelihood) method which we recommend. The best fit value of the EC50 (multiplied by \(10^6\) to convert to micromolar) is 2.4 \(\mu M\) with the 95% confidence interval ranging from 0.85 \(\mu M\) to 15.3 \(\mu M\).

Prism also reports these values on a log scale. The logEC50 (log molar) is -5.62 with a 95% confidence interval ranging from -6.07 to -4.82.

Three things to notice:

- The confidence interval is fairly wide. This is because the data have some scatter and also because we are fitting the EC50 plus three other parameters (the top and bottom plateaus and the slope). If you fix one or more of those parameters to constant values, the confidence interval for the EC50 would get narrower.

- The confidence interval, expressed on a concentration scale, is quite asymmetrical.

- The confidence interval, expressed on a log scale, is close to symmetrical.

An alternative way to fit the data are to transform the X values to their logarithm, and then fit the results of this transform to the model: log[Agonist] vs. response -- variable slope with the Confidence tab set to compute confidence intervals using the asymmetrical (likelihood) method which we recommend using always. You'll get exactly the same results this way.

**Symmetrical confidence intervals**

Prism 7 offers two methods to compute the confidence intervals. The results above were computed using the new method that produces asymmetrical intervals. If you use other method (the only method offered in prior versions of Prism), the confidence intervals for the parameter you fit are always symmetrical.

If you fit the model that fits the logEC50, the confidence interval ranges from -6.07 to -4.82. The "asymmetrical" confidence interval of the logEC50 shown above is nearly symmetrical, so the symmetrical confidence interval is nearly the same (is the same to three digits).
But what if you fit the EC50 and ask for symmetrical confidence interval. In this case the reported interval ranges from 0 \( \mu \text{M} \) to 5.21\( \mu \text{M} \). This seems asymmetrical because the EC50 is constrained to be greater than zero. If you removed that constraint, Prism would report a confidence interval ranging from -0.42 \( \mu \text{M} \) to 5.21\( \mu \text{M} \), which is symmetrical around the best-fit value of 2.4 \( \mu \text{M} \). This symmetrical interval does a really poor job of quantifying the precision of the IC50. The simple fact is that the uncertainty is asymmetrical, and a symmetrical interval can be very misleading. Even setting aside the negative portion of the confidence interval, it includes all values from zero on up, so doesn't provide any guidance about how low the EC50 might really be. That is why we recommend that you choose asymmetrical intervals routinely.

**Do not transform the standard error of the logEC50**

We suggest that you always ask Prism to report the asymmetrical confidence intervals, and not to report the standard error of the parameters. The SE of the parameters does a poor job of quantifying precision. It gets particularly confusing when logarithms get involved. When some people see the SE of the logEC50, they are tempted to convert this to the standard error of the EC50 by taking the antilog. This is invalid.

In the example, the SE of the logEC50 is 0.2158. The antilog of 0.2158 equals 1.644. This value is not the SE of the EC50. It cannot be interpreted as a plus/minus error value. It is a times/divided by error value. To get an equivalent to the logEC50 plus or minus one of the logEC50, find the lower limit by dividing the EC50 by 1.644 and the upper limit by multiplying by 1.644. But this is not commonly done, so it would be confusing if you tried to report results this way.

**Averaging the EC50 from several experiments**

The uncertainty is close to symmetrical when you express the midpoint of a dose-response curve as a logEC50, but is usually far from symmetrical when you express it as the EC50. When pooling several experiments, therefore, it is best to average the logEC50 values, which will give a different result than averaging the EC50 values.
3.6.1.6 Hill slope

The usual steepness of a dose-response curve

Many dose-response curves follow the shape of a receptor binding curve. As shown below, 81 times more agonist is needed to achieve 90% response than a 10% response.

Since the linkage between agonist binding and response can be very complex, any shape is possible. It seems surprising, therefore, that so many dose-response curves have shapes almost identical to receptor binding curves, even when we know there are multiple steps between binding and measured response. It turns out that no matter how many steps intervene between agonist binding and response, the dose-response curve will have the usual steepness so long as each messenger binds to a single binding site according to the law of mass action.

The slope factor or Hill slope

Some dose-response curves are steeper or shallower than the standard curve. The steepness is quantified by the Hill slope, also called a slope factor. A dose-response curve with a standard slope has a Hill slope of 1.0. A steeper curve has a higher slope factor, and a shallower curve has a lower slope factor.
If you use a single concentration of agonist and varying concentrations of antagonist, the curve goes downhill and the slope factor is negative. The steeper the downhill slope, the more negative the Hill slope.

**Standard slope or variable slope?**

Because this standard slope is so common, Prism comes with equations with the standard slope built in. The equations that don't have 'variable slope' in their name assume the standard slope (1.0 for stimulation, -1.0 for inhibition).

Deciding whether to fit a model with a standard slope or a variable slope is not easy.

If you have lots of data points (more than a dozen, perhaps lots more), then you can fit the slope by picking a variable slope equation. If you have fewer data points, and a standard system, it makes sense to choose an equation with a standard slope.

### 3.6.1.7 Choosing a dose-response equation

To choose a dose-response model in Prism, you need to answer four questions:
Are the X values concentrations (or doses) or the logarithm of concentration or dose?

Prism 7 offers two sets of equations for dose-response curves. One set assumes X is the logarithm of dose or concentration. The other set assumes X is the dose or concentration. Be sure to pick an equation that matches the data.

Stimulation or inhibition?

Prism offers one set of dose-response equations for stimulation and another set for inhibition. The inhibitory equations are set up to run downhill. The only difference is that the inhibitory equations fit the IC50 ("I" for inhibition) while the stimulation equations fit the EC50 ("E" for effective).

If the curve goes up hill, choose from the set of stimulation equations. If the curve goes down hill, choose from the set of inhibition equations.

Standard slope or variable slope?

Prism offers equations using a standard slope, which have a Hill slope of 1.00 (for stimulation) or -1.00 (for inhibition), and variable slope (fit by Prism). The equations that don't have 'variable slope' in their name assume the standard slope. The standard slope is expected when measuring binding of a ligand to a receptor where there is no heterogeneity or cooperativity. But it turns out that many other log(dose) vs. response curves have the same standard slope.

If you have lots of data points (more than a dozen, perhaps lots more), then you can fit the slope by picking a variable slope equation. If you have fewer data points, and a standard system, it makes sense to choose an equation with a standard slope.

There are many situations where it is not clear which approach is better.

Normalized or not?

If your data have been normalized so the curve runs from Y= 0 to Y=100, you may wish to choose a normalized model. These models don't fit the bottom and top plateaus, but rather force the bottom plateau to equal 0 and the top plateau to equal 100. Only choose a 'normalized response' equation when you have determined the values that define 0
and 100 very precisely. Just because the data have been normalized doesn't mean to have to constrain the curve in that way. \[^{278}\]

There are many situations where it is not clear whether it makes sense to use a normalized model or not. It is not always a straightforward decision.

**Special dose-response models**

Prism has a set of special models used for special dose-response situations:

- **Asymmetrical (five parameter)** \[^{279}\]
- **Biphasic dose-response** \[^{280}\]
- **Bell-shaped dose-response** \[^{281}\]
- **Operational model - Depletion** \[^{282}\]
- **Operational model - Partial agonist** \[^{283}\]
- **Gaddum/Schild EC50 shift** \[^{284}\]
- **Allosteric EC50 shift** \[^{285}\]
- **ECanything** \[^{286}\]

**3.6.1.8 Pros and cons of normalizing the data**

The dose-response model has four parameters: the bottom plateau, the top plateau, the EC50, and the slope factor (which is often constrained to a standard value).

The main goal of fitting the dose-response curve in many situations is to determine the best-fit value of the EC50, which is the concentration that provokes a response halfway between the top and bottom plateaus. If those plateaus are not well defined, the EC50 will be very uncertain. Think of it this way: If you have not defined "100" and "0" very precisely, you also have not defined "50" precisely, and therefore cannot determine the EC50 precisely.
One way to solve the problem is to constrain the Top or Bottom, or both, to control values. Another alternative is to normalize your data so responses run from 0 to 100, and then choose a "normalized response" model. These models don't fit the bottom and top plateaus, but rather force the bottom plateau to equal 0 and the top plateau to equal 100. Only choose a 'normalized response' equation when you have determined the values that define 0 and 100 very precisely.

Prism makes it easy to normalize the data so the values run from 0% to 100%. Simply click Analyze, choose the Normalize analysis., and define how 0% and 100% are defined. When fitting a dose-response curve, you can fit either the raw data or normalized data.

Notes:

- It is not necessary to normalize before fitting dose-response data. In many cases, it is better to show the actual data.

- You can only plot several different dose-response curves on one graph using one axis when they are comparable. If the different experiments measured different variables, normalizing puts them into comparable units. This can be useful.

- Whether or not you choose to normalize your data, you still need to choose how to fit the data. Do you want Prism to find best-fit values for the Top and Bottom plateaus? Or do you want those plateaus to be determined by control data?

- If you normalize your data, you can choose one of the normalized dose-response equations. These constrain the the curve to run from 0% to 100%. This kind of constraint only makes sense, when 0% and 100% are defined by good control data. If the definitions of 0% and 100% are ambiguous, then so is the definition of "50%", and thus the EC50 is also ambiguous.

- Just because you chose to normalize your data doesn't mean you must constrain the curve to run from 0 to 100%. You may prefer to have Prism fit those two plateaus.

- If you don't normalize your data, you can use the Constrain tab to fix Top and Bottom to values determined from control experiments. So the decision to constrain Top and Bottom is quite distinct from the decision to normalize your data before fitting.
• It is possible, and can be reasonable, to fix one of those parameters (Top or Bottom) to a constant value but not the other.

• If you normalize, **don't also choose to differentially weight the data**.

3.6.1.9 The term "logistic"

The term *logistic* has three meanings which have little relationship to each other (1).

**Meaning 1: Logistic population growth**

The term "logistic" was first invented in the nineteenth century to describe population growth curves. The idea is pretty simple. Population growth is limited, so can't ever exceed some value we'll call Nmax. The rate of change of population at any time t is proportional to the number of individuals alive at that time (Nt). But population growth slows down as it reaches the maximum, so is also proportional to (Nmax - Nt). So the rate of change of population is proportional to Nt(Nmax - Nt).

**Integrate that differential equation**, and the result is called a logistic equation. It defines a sigmoidal shaped curve that defines the population at any time. The model has three parameters: the starting population, the maximum population, and the time it takes to reach half-maximal. Sometimes it is modified to add a fourth parameter to define the steepness of the curve.

This curve was used by demographers in the past, but actually doesn't do a very good job of describing the growth of human populations. It is still used to model the growth of tumors, and to model the fraction of a population that uses a new product (like a mobile phone).
This model is also used for autocatalytic reactions, where the product of the reaction is also a catalyst for that reaction. With this kind of reaction, the rate of product accumulation is proportional to the concentration of product already produced times the concentration of remaining substrate. This has the same mathematical form as the population growth model. The graph is identical to the one above, except the Y-axis would be the concentration of the product produced by the enzyme reaction (instead of population).

Meaning 2: "Logistic" dose-response curves

Bindslev reviews the history of the many equations used to describe dose response curves(2). A common equation, based on work by Hill, defines a response as a function of the minimum response, the maximum response, the concentration required to evoke a response half-way between the minimum and maximum (the EC50), and a parameter that describes how steep the curve is. In Prism, this equation is called the log(agonist) vs response curve -- Variable slope (four parameters).

Response at any dose is defined by an equation with four parameters: minimum response, maximum response, the log(EC50), and the slope factor (also called the Hill slope). Surprisingly, this equation has the same mathematical form as the logistic equation describing population growth, even though the two uses are entirely different. For this reason, the dose-response equation is sometimes called a "logistic" equation. Since there are four parameters, it is sometimes called the four parameter logistic equation, abbreviated 4PL.

This dose-response model describes many systems, but it is purely empirical. It does not correspond to any molecular model (except, perhaps, when the Hill slope is an integer).
Note the huge differences between the use of a logistic equation to describe population growth, and its use to describe dose-response curves.

- The X axis of the population growth curve is time. The X axis of the dose-response curve is the logarithm of dose or concentration.

- The equation for population growth comes from theory. The equation for dose-response relationships is empirical.

- Despite its name, no logarithms are used in the logistic equation for population growth. The X axis of the logistic dose-response curve is the logarithm of dose or concentration.

**Meaning 3: Logistic regression**

The third use of the word *logistic*, completely unrelated to the other two uses, is logistic regression. This is a multiple regression method used when the outcome (dependent variable) has two possible outcomes. The point of the equation is to predict the probability, \( P \), of an outcome from a set of independent variables. To create a model that always predicts a value for \( P \) that is between 0 and 1 the logistic regression model actually predicts the natural logarithm of the odds. The function that computes the natural logarithm of the odds from a fraction is called the *logit* function (pronounced with a long O and a soft G), so regression used to predict the logit of a probability from multiple independent variables is called *logistic regression*.

**References**


3.6.1.10 50% of what? Relative vs absolute IC50

IC50 can be ambiguous

50% of what? There can be more than one answer in some situations.

The concepts of IC50 is fundamental to pharmacology. The IC50 is the concentration of an inhibitor where the response (or binding) is reduced by half. Seems simple enough. But when you actually go to fit data to determine these values, there are several complexities and ambiguities.

The rest of this article is about IC50 (I for inhibition, for downward sloping dose-response curves). All the ideas can be applied to stimulatory curves and EC50 (E for effective) as well. Just stand on your head when you view the figures!.

The ideal situation

This figure shows an ideal situation:

The green symbols show measurements made with controls. The ones on the left (Blank) have no inhibitor, so define "100%". The ones on the right are in the presence of a maximal concentration of a standard inhibitor, so define "0%". The data of the experimental dose-response curve (red dots) extend all the way between the two control values.
When fitting this curve, you need to decide how to fit the top plateau of the curve. You have three choices:

- Fit the data only, ignoring the Blank control values.

- Average the Blank control values, and set the parameter Top to be a constant value equal to the mean of the blanks.

- Enter the blank values as if they were part of the dose-response curve. Simply enter a low dose, perhaps 10-10 or 10-11. You can't enter zero, because zero is not defined on a log scale.

The results will be very similar with any of these methods, because the data form a complete dose-response curve with a clear top plateau that is indistinguishable from the blank. I prefer the third method, as it analyzes all the data, but that is not a strong preference.

Similarly, there are three ways to deal with the bottom plateau: Fit the data only, set Bottom to be a constant equal to the average of the NS controls, and put the NS controls into the fit as if they were a very high concentration of inhibitor.

That is the ideal situation. There is no ambiguity about what IC50 means.

**A potentially ambiguous situation**

The graph below shows an unusual situation where the inhibition curve plateaus well above the control values (NS) defined by a high concentration of a standard drug. This leads to alternative definitions of IC50.
Clearly, a single value cannot summarize such a curve. You'd need at least two values, one to quantify the middle of the curve (the drug's potency) and one to quantify how low it gets (the drug's maximum effect).

The graph above shows two definitions of the IC50.

The **relative IC50** is by far the most common definition, and the adjective *relative* is usually omitted. It is the concentration required to bring the curve down to point half way between the top and bottom plateaus of the curve. The NS values are totally ignored with this definition of IC50. This definition is the one upon which classical pharmacological analysis of agonist and antagonist interactions is based. With appropriate consideration of the biological system and concentrations of interacting ligands, estimated Kd values can often be derived from the IC50 value defined this way (not so for the "so-called absolute IC50" mentioned below).

The concentration that provokes a response halfway between the Blank and the NS value is sometimes called the **absolute IC50**, The horizontal dotted lines show how 100% and 0% are defined, which then defines 50%. This term is not entirely standard. Since this value does not quantify the potency of a drug, the authors of the International Union of Pharmacology Committee on Receptor Nomenclature (1) think that...
the concept of absolute IC50 (and that term) is not useful (R. Neubig, personal communication). I agree.

The concept (but not the term "absolute IC50") is used to quantify drugs that slow cell growth. The abbreviation GI50 is used for what we call here the absolute IC50. It is also used by the Environmental Protection Agency (EPA) in evaluating endocrine disrupters (Appendix A). That document uses the term IC50 to refer to the absolute IC50, and the term EC50 to refer to the relative IC50. It doesn't use the terms relative and absolute.

If you really want to use the absolute IC50, the next page has instructions for fitting a curve to find it.

Reference


Download the Prism file used to create all the graphs in this article.

3.6.1.11 Fitting the absolute IC50

Fitting a dose-response curve to find the absolute IC50

The concept of an absolute IC50 is not standard, and many find it not to be useful. But if you do, it is not hard to fit a curve to determine it. This EPA document, give the needed equation (which I have generalized a bit, so not require that the data already be normalized).

\[ \text{Fifty} = \frac{(\text{Top} + \text{Baseline})}{2} \]

\[ Y = \text{Bottom} + \frac{(\text{Top} - \text{Bottom})}{(1+10^{(\text{LogIC50\,-\,X}\times\text{HillSlope})})} + \log\left(\frac{\text{Top} - \text{Bottom}}{\text{Fifty} - \text{Bottom}}\right) \]

Note the distinction between the parameter Bottom and Baseline. Bottom is the Y value of the bottom plateau of the curve itself. Baseline is the Y value that defines 0% -- maximal inhibition by a standard drug. You'll definitely want to constrain Baseline to be a constant value based on controls. You may also want to constrain Top.

Download the Prism file that fits that equation to make the graph shown above. When fitting data to that equation, don't forget to constrain
Baseline and Top to appropriate values determined by controls. Additionally, this file contains another graph where the data are already normalized to run from 0 to 100%. These data are fit to a simpler equation where Baseline is set to equal zero, and Top is set to equal 100. These are hard wired into the equation, so you don't have to remember to constrain those two parameters to constant values.

**An alternative approach for normalized data**

Here is an alternative approach you can use if your data are normalized. It does not require entering a user-defined equation.

1. Make sure that your data are normalized to some controls. That means the response at the left end (low concentrations) is near 100%, and the response at higher concentrations is above 0%. If the response plateaus at 0%, then an absolute and relative IC50 are the same, and you can just fit the usual dose-response curve to find the IC50.

2. At the bottom of the data table, add a new row of data. Enter 50 into each Y column. Leave X blank for this row.

3. Use nonlinear regression to fit the data to the log(inhibitor) vs. response (variable slope) curve.

4. On the first (fit) tab of the nonlinear regression dialog, check the option: "Interpolate unknowns from standard curve." Of course, these data are not a standard curve, and there are unknowns. But this option asks Prism to interpolate the X value of the curve when Y=50.

5. On the Constrain tab, consider constraining Top to have a constant value of 100. If you have used good controls to normalize your data, then you know the top plateau of the curve has to be Y=100, so should tell Prism to use that constraint.

6. The IC50 reported as part of the main results table will be the relative IC50.

7. Look on the additional results page of interpolated results to see the value of X when Y=50. This is the absolute IC50.
3.6.1.12 Incomplete dose-response curves

Any attempt to determine an IC50 by fitting a curve to the data in the graph above will be useless. Prism might, or might not, be able to fit a dose-response curve to the data. But if the curve fits, the value of the IC50 is likely to be meaningless and have a very wide confidence interval. The data simply don't really define a top plateau (which would define 100) and certainly doesn't provide even a hint of a bottom plateau (which would define 0). If data haven't defined 100 or 0, then 50 is undefined too, as is the IC50.

If you also have control values that define 100 and 0, then the curve can be easily fit. The curve below was created by fitting a dose response curve, but constraining the Top plateau to be a constant value equal to the mean of the Blanks values, and the Bottom plateau equal to the mean of the NS values. Note that the blank and NS values are shown in green.
The value of the IC50 fit this way only makes sense if you assume that higher concentrations of the inhibitor would eventually inhibit down to the NS values. That is an assumption that can't be tested with the data at hand.

The distinction between relative and absolute IC50 doesn't really apply to these data. Because the data don't define a bottom plateau, the IC50 can only be defined relative to the NS control values.

### 3.6.1.13 Troubleshooting fits of dose-response curves

**First consult the analysis checklists**

If your goal is to interpolate from the standard curve, look at the analysis checklist designed for that purpose.

If your goal is to fit a model to determine parameter values, review the general analysis checklist for fitting a model.

**Questions to ask about dose-response curves**

- Are the X values concentrations (or doses) or the logarithm of concentrations (or doses)? Prism offers equations for each. You can also use the Transform analysis to make a new results table where the X values are logarithms. Note that stretching an axis to a log scale is not at all the same thing. Prism's nonlinear regression fits the values it is given. Using the Format Axis dialog to stretch the X axis
to a logarithmic scale does not change the values the curve fitter sees.

- Is the top plateau defined by the data? If not, can it be constrained or shared? If the top plateau is not defined either by the data or by a constraint, then any fit of a dose-response curve is unlikely to be useful.

- Is the bottom plateau defined by the data? If not, can it be constrained or shared? If the bottom plateau is not defined either by the data or by a constraint, then any fit of a dose-response curve is unlikely to be useful.

- Were the data normalized? To what values? If the data were normalized, consider constraining the top and bottom plateaus to be 0 and 1 (or 100) in the Constraint tab?

- Is the Hill slope fixed? A Hill slope of 1.0 or -1.0 is commonly seen in many systems, but not all.

- Four parameter assumes symmetry. The usual equations to fit dose response curves have four parameters (top, bottom, EC50, Hill Slope), and define symmetrical curves. You can choose an equation that adds a fifth parameter to fit asymmetrical curves.

- Are you fitting the relative IC50 but expecting an absolute IC50, and so are surprised by the results.

3.6.2 Dose-response - Stimulation
3.6.2.1 Equation: log(agonist) vs. response

This equation is used when X values are logarithms of doses or concentrations. Use a related equation when X values are concentrations or doses.

**Introduction**

Many log(dose) vs. response curves follow the familiar symmetrical sigmoidal shape. The goal is to determine the EC50 of the agonist - the concentration that provokes a response half way between the basal (Bottom) response and the maximal (Top) response.
This model assumes that the dose response curve has a **standard slope**, equal to a Hill slope (or slope factor) of 1.0. This is the slope expected when a ligand binds to a receptor following the law of mass action, and is the slope expected of a dose-response curve when the second messenger created by receptor stimulation binds to its receptor by the law of mass action. If you don't have many data points, consider using the standard slope model. If you have lots of data points, pick the variable slope model to determine the Hill slope from the data.

This equation is sometimes called a three parameter dose-response curve. If you also fit the Hill slope, then it is a four parameter equation.

**Step by step**

Create an XY data table. Enter the logarithm of the concentration of the agonist into X. Enter response into Y in any convenient units. Enter one data set into column A, and use columns B, C... for different treatments, if needed.

If you prefer to enter concentrations, rather than the logarithm of concentrations, use Prism to transform the X values to logs."

From the data table, click Analyze, choose nonlinear regression, choose the panel of equations "Dose-response curves - Stimulation" and then choose the equation \( \log(\text{Agonist}) \text{ vs. response} \).

If you have subtracted off any basal response, consider constraining Bottom to a constant value of 0.

**Model**

\[
Y = \text{Bottom} + \frac{(\text{Top}-\text{Bottom})}{(1+10^{((\text{LogEC50}-X))})}
\]
Interpret the parameters

**EC50** is the concentration of agonist that gives a response half way between Bottom and Top. This is not the same as the response at Y=50. Depending on which units Y is expressed in, and the values of Bottom and Top, the EC50 may give a response nowhere near "50". Prism reports both the EC50 and its log.

**Top** and **Bottom** are plateaus in the units of the Y axis.

3.6.2.2 Equation: log(agonist) vs. response -- Variable slope

This equation is used when X values are logarithms of doses or concentrations. Use a related equation when X values are concentrations or doses.

Introduction

Many log(dose) response curves follow the familiar symmetrical sigmoidal shape. The goal is to determine the EC50 of the agonist - the concentration that provokes a response half way between the basal (Bottom) response and the maximal (Top) response.

Many dose-response curves have a standard slope of 1.0. This model does not assume a standard slope but rather fits the Hill Slope from the data, and so is called a **Variable slope** model. This is preferable when
you have plenty of data points. It is also called a four-parameter dose-response curve, or four-parameter logistic curve, abbreviated 4PL.

**Step by step**

Create an XY data table. Enter the logarithm of the concentration of the agonist into X. Enter response into Y in any convenient units. Enter one data set into column A, and use columns B, C... for different treatments, if needed.

If you prefer to enter concentrations, rather than the logarithm of concentrations, use Prism to transform the X values to logs.

From the data table, click Analyze, choose nonlinear regression, choose the panel of equations "Dose-response curves - Stimulation" and then choose the equation "log(Agonist) vs. response -- Variable slope".

Consider constraining the parameter HillSlope to its standard values of 1.0. This is especially useful if you don't have many data points, and therefore cannot fit the slope very well.

If you have subtracted off any basal response, consider constraining Bottom to a constant value of 0.

**Model**

\[ Y = \text{Bottom} + \frac{(\text{Top} - \text{Bottom})}{(1 + 10^{((\text{LogEC50} - X) \times \text{HillSlope})})} \]
Interpret the parameters

**EC50** is the concentration of agonist that gives a response half way between Bottom and Top. This is not the same as the response at Y=50. Depending on which units Y is expressed in, and the values of Bottom and Top, the EC50 may give a response nowhere near "50". Prism reports both the EC50 and its log.

**HillSlope** describes the steepness of the family of curves. A HillSlope of 1.0 is standard, and you should consider constraining the Hill Slope to a constant value of 1.0. A Hill slope greater than 1.0 is steeper, and a Hill slope less than 1.0 is shallower.

**Top** and **Bottom** are plateaus in the units of the Y axis.

3.6.2.3 Equation: log(agonist) vs. normalized response

This equation is used when X values are logarithms of doses or concentrations. Use a related equation when X values are concentrations or doses.

**Introduction**

Many log(dose) vs. response curves follow the familiar symmetrical sigmoidal shape.

If you have good control data, it can make sense to normalize the response to run between 0% and 100%. This model assumes that the data have been normalized, so forces the curve to run from 0% to 100%. The goal is to determine the EC50 of the agonist - the concentration that provokes a response equal to 50%.

It only makes sense to fit a normalized model when you are sure you have defined 0% and 100% quite accurately. If your data define a complete sigmoidal curve, it is best to fit the entire curve and let Prism fit the Top and Bottom plateaus. If your data don't form a full sigmoidal curve, but you can define the bottom and top by solid control data, then fitting to a normalized model is preferable.

This model assumes that the dose response curve has a **standard slope**, equal to a Hill slope (or slope factor) of 1.0. This is the slope.
expected when a ligand binds to a receptor following the law of mass action, and is the slope expected of a dose-response curve when the second (and third...) messengers created by receptor stimulation binds to its receptor by the law of mass action. If you don't have many data points, consider using the standard slope model. If you have lots of data points, pick the variable slope model to determine the Hill slope from the data.

**Step by step**

Create an XY data table. Enter the logarithm of the concentration of the agonist into X. Enter response into Y in any convenient units. Enter one data set into column A, and use columns B, C... for different treatments, if needed.

If you prefer to enter concentrations, rather than the logarithm of concentrations, use Prism to transform the X values to logs.[269]

From the data table, click Analyze, choose nonlinear regression, choose the panel of equations "Dose-response curves - Stimulation" and then choose the equation "log(Agonist) vs. normalized response".

**Model**

\[ Y = \frac{100}{1 + 10^{(\log EC50 - X)}} \]
In **Interpret the parameter**

**EC50** is the concentration of agonist that gives a response halfway between Bottom and Top. This is not the same as the response at Y=50. Depending on which units Y is expressed in, and the values of Bottom and Top, the EC50 may give a response nowhere near "50". Prism reports both the EC50 and its log.

### 3.6.2.4 Equation: log(agonist) vs. normalized response -- Variable slope

This equation is used when X values are logarithms of doses or concentrations. Use a related equation when X values are concentrations or doses.

**Introduction**

Many log(dose) response curves follow the familiar symmetrical sigmoidal shape.

If you have good control data, it can make sense to normalize the response to run between 0% and 100%. This model assumes that the data have been normalized, so forces the curve to run from 0% to 100%. The goal is to determine the EC50 of the agonist - the concentration that provokes a response equal to 50%.

It only makes sense to fit a normalized model when you are sure you have defined 0% and 100% quite accurately. If your data define a complete sigmoidal curve, it is best to fit the entire curve and let Prism fit the Top and Bottom plateaus. If your data don't form a full sigmoidal curve, but you can define the bottom and top by solid control data, then fitting to a normalized model is preferable.

Many dose-response curves have a standard slope of 1.0. This model does not assume a standard slope but rather fits the Hill Slope from the data, and so is called a **Variable slope** model. This is preferable when you have plenty of data points.

**Step by step**

Create an XY data table. Enter the logarithm of the concentration of the agonist into X. Enter response into Y in any convenient units. Enter one
data set into column A, and use columns B, C... for different treatments, if needed.

If you prefer to enter concentrations, rather than the logarithm of concentrations, use Prism to transform the X values to logs.

From the data table, click Analyze, choose nonlinear regression, choose the panel of equations "Dose-response curves - Stimulation" and then choose the equation "log(Agonist) vs. normalized response -- Variable slope".

**Model**

\[ Y = \frac{100}{1 + 10^{((\log EC50 - X) * \text{HillSlope})}} \]

**Interpret the parameters**

**EC50** is the concentration of agonist that gives a response half way between Bottom and Top. This is not the same as the response at \( Y = 50 \). Depending on which units \( Y \) is expressed in, and the values of Bottom and Top, the EC50 may give a response nowhere near "50". Prism reports both the EC50 and its log.

**HillSlope** describes the steepness of the family of curves. A HillSlope of 1.0 is standard, and you should consider constraining the Hill Slope to a constant value of 1.0.
3.6.2.5 Equation: [Agonist] vs. response

This equation is used when X values are concentrations. Use a related equation when X values are logarithms of concentrations or doses.

Introduction

Many log(dose) vs. response curves follow the familiar symmetrical sigmoidal shape. The goal is to determine the EC50 of the agonist - the concentration that provokes a response half way between the basal (Bottom) response and the maximal (Top) response.

This model assumes that the dose response curve has a standard slope, equal to a Hill slope (or slope factor) of 1.0. This is the slope expected when a ligand binds to a receptor following the law of mass action, and is the slope expected of a dose-response curve when the second messenger created by receptor stimulation binds to its receptor by the law of mass action. If you don't have many data points, consider using the standard slope model. If you have lots of data points, pick the variable slope model to determine the Hill slope from the data.

This equation is sometimes called a three parameter dose-response curve. If you also fit the Hill slope, then it is a four parameter equation.

Step by step

Create an XY data table. Enter the concentration of the agonist into X. Enter response into Y in any convenient units. Enter one data set into column A, and use columns B, C... for different treatments, if needed.

From the data table, click Analyze, choose nonlinear regression, choose the panel of equations "Dose-response curves - Stimulation" and then choose the equation [Agonist] vs. response.

If you have subtracted off any basal response, consider constraining Bottom to a constant value of 0.

Since the uncertainty of the EC50 is very asymmetric, be sure to choose to compute the confidence intervals using the likelihood ratio asymmetric method.

Double click on the X axis of the graph, and choose (at the upper left of the Format Graph dialog) to stretch the axis to a logarithmic scale.
**Model**

\[ Y = \text{Bottom} + X \times \frac{\text{Top} - \text{Bottom}}{\text{EC50} + X} \]

**Interpret the parameters**

- **EC50** is the concentration of agonist that gives a response half way between Bottom and Top. This is not the same as the response at \( Y = 50 \). Depending on which units \( Y \) is expressed in, and the values of Bottom and Top, the EC50 may give a response nowhere near "50". Prism reports both the EC50 and its logarithm.

- **Top** and **Bottom** are plateaus in the units of the Y axis.

**Introduction**

Many log(dose) response curves follow the familiar symmetrical sigmoidal shape. The goal is to determine the EC50 of the agonist - the concentration that provokes a response half way between the basal (Bottom) response and the maximal (Top) response.
Many dose-response curves have a standard slope of 1.0. This model does not assume a standard slope but rather fits the Hill Slope from the data, and so is called a **Variable slope** model. This is preferable when you have plenty of data points. It is also called a four-parameter dose-response curve, or four-parameter logistic curve, abbreviated 4PL.

**Step by step**

Create an XY data table. Enter the concentrations of the agonist into X. Enter response into Y in any convenient units. Enter one data set into column A, and use columns B, C... for different treatments, if needed.

From the data table, click Analyze, choose nonlinear regression, choose the panel of equations "Dose-response curves - Stimulation" and then choose the equation "[Agonist] vs. response -- Variable slope".

Consider constraining the parameter HillSlope to its standard values of 1.0. This is especially useful if you don't have many data points, and therefore cannot fit the slope very well.

If you have subtracted off any basal response, consider constraining Bottom to a constant value of 0.

Since the uncertainty of the EC50 is very asymmetric, be sure to choose to compute the confidence intervals using the **likelihood ratio asymmetric method**.

Double click on the X axis of the graph, and choose (at the upper left of the Format Graph dialog) to stretch the axis to a logarithmic scale.

**Model**

\[
Y = Bottom + (X^{HillSlope} \times (Top-Bottom))/(X^{HillSlope} + EC50^{HillSlope})
\]
Interpret the parameters

**EC50** is the concentration of agonist that gives a response half way between Bottom and Top. This is not the same as the response at Y=50. Depending on which units Y is expressed in, and the values of Bottom and Top, the EC50 may give a response nowhere near "50". Prism reports both the EC50 and its log.

**HillSlope** describes the steepness of the family of curves. A HillSlope of 1.0 is standard, and you should consider constraining the Hill Slope to a constant value of 1.0. A Hill slope greater than 1.0 is steeper, and a Hill slope less than 1.0 is shallower.

**Top** and **Bottom** are plateaus in the units of the Y axis.

### 3.6.2.7 Equation: [Agonist] vs. normalized response

This equation is used when X values are concentrations. Use a related equation when X values are logarithms of concentrations or doses.

**Introduction**

Many log(dose) vs. response curves follow the familiar symmetrical sigmoidal shape.
If you have good control data, it can make sense to normalize the response to run between 0% and 100%. This model assumes that the data have been normalized, so forces the curve to run from 0% to 100%. The goal is to determine the EC50 of the agonist - the concentration that provokes a response equal to 50%.

It only makes sense to fit a normalized model when you are sure you have defined 0% and 100% quite accurately. If your data define a complete sigmoidal curve, it is best to fit the entire curve and let Prism fit the Top and Bottom plateaus. If your data don't form a full sigmoidal curve, but you can define the bottom and top by solid control data, then fitting to a normalized model is preferable.

This model assumes that the dose response curve has a standard slope, equal to a Hill slope (or slope factor) of 1.0. This is the slope expected when a ligand binds to a receptor following the law of mass action, and is the slope expected of a dose-response curve when the second (and third...) messengers created by receptor stimulation binds to its receptor by the law of mass action. If you don't have many data points, consider using the standard slope model. If you have lots of data points, pick the variable slope model to determine the Hill slope from the data.

**Step by step**

Create an XY data table. Enter the concentrations of the agonist into X. Enter response into Y in any convenient units. Enter one data set into column A, and use columns B, C... for different treatments, if needed.

From the data table, click Analyze, choose nonlinear regression, choose the panel of equations "Dose-response curves - Stimulation" and then choose the equation "[Agonist] vs. normalized response".

Since the uncertainty of the EC50 is very asymmetric, be sure to choose to compute the confidence intervals using the likelihood ratio asymmetric method.

Double click on the X axis of the graph, and choose (at the upper left of the Format Graph dialog) to stretch the axis to a logarithmic scale.

**Model**

\[ Y = \frac{100 \times X}{EC50 + X} \]
Interpret the parameter

**EC50** is the concentration of agonist that gives a response half way between Bottom and Top. This is not the same as the response at Y=50. Depending on which units Y is expressed in, and the values of Bottom and Top, the EC50 may give a response nowhere near "50". Prism reports both the EC50 and its log.

3.6.2.8 Equation: [Agonist] vs. normalized response -- Variable slope

This equation is used when X values are concentrations. Use a related equation when X values are logarithms of concentrations or doses.

Introduction

Many log(dose) response curves follow the familiar symmetrical sigmoidal shape.

If you have good control data, it can make sense to normalize the response to run between 0% and 100%. This model assumes that the data have been normalized, so forces the curve to run from 0% to 100%. The goal is to determine the EC50 of the agonist - the concentration that provokes a response equal to 50%.
It only makes sense to fit a normalized model when you are sure you have defined 0% and 100% quite accurately. If your data define a complete sigmoidal curve, it is best to fit the entire curve and let Prism fit the Top and Bottom plateaus. If your data don't form a full sigmoidal curve, but you can define the bottom and top by solid control data, then fitting to a normalized model is preferable.

Many dose-response curves have a standard slope of 1.0. This model does not assume a standard slope but rather fits the Hill Slope from the data, and so is called a Variable slope model. This is preferable when you have plenty of data points.

**Step by step**

Create an XY data table. Enter the concentrations of the agonist into X. Enter response into Y in any convenient units. Enter one data set into column A, and use columns B, C... for different treatments, if needed.

From the data table, click Analyze, choose nonlinear regression, choose the panel of equations "Dose-response curves - Stimulation" and then choose the equation "[Agonist] vs. normalized response -- Variable slope".

Since the uncertainty of the EC50 is very asymmetric, be sure to choose to compute the confidence intervals using the likelihood ratio asymmetric method.

Double click on the X axis of the graph, and choose (at the upper left of the Format Graph dialog) to stretch the axis to a logarithmic scale.

**Model**

\[ Y = 100 \times (X^{\text{HillSlope}})/(EC50^{\text{HillSlope}} + (X^{\text{HillSlope}})) \]
Interpret the parameters

**EC50** is the concentration of agonist that gives a response half way between Bottom and Top. This is not the same as the response at Y=50. Depending on which units Y is expressed in, and the values of Bottom and Top, the EC50 may give a response nowhere near "50". Prism reports both the EC50 and its log.

**HillSlope** describes the steepness of the family of curves. A HillSlope of 1.0 is standard, and you should consider constraining the Hill Slope to a constant value of 1.0.

3.6.3 Dose-response - Inhibition

3.6.3.1 Equation: log(inhibitor) vs. response

This equation is used when X values are logarithms of doses or concentrations. Use a related equation when X values are concentrations or doses.

Introduction

Many log(inhibitor) vs. response curves follow the familiar symmetrical sigmoidal shape. The goal is to determine the IC50 of the inhibitor - the concentration that provokes a response half way between the maximal (Top) response and the maximally inhibited (Bottom) response.
This model assumes that the dose response curves has a **standard slope**, equal to a Hill slope (or slope factor) of -1.0. This is the slope expected when a ligand binds to a receptor following the law of mass action, and is the slope expected of a dose-response curve when the second messenger created by receptor stimulation binds to its receptor by the law of mass action. If you don't have many data points, consider using the standard slope model. If you have lots of data points, pick the variable slope model to determine the Hill slope from the data.

**Step by step**

Create an XY data table. Enter the logarithm of the concentration of the inhibitor into X. Enter response into Y in any convenient units. Enter one data set into column A, and use columns B, C... for different treatments, if needed.

If you prefer to enter concentrations, rather than the logarithm of concentrations, use Prism to transform the X values to logs.

From the data table, click Analyze, choose nonlinear regression, choose the panel of equations "Dose-response curves - Inhibition" and then choose the equation "log(inhibitor) vs. response".

If you have subtracted off any basal response, consider constraining Bottom to a constant value of 0.

**Model**

\[ Y = \text{Bottom} + \frac{(\text{Top}-\text{Bottom})}{(1+10^{(X-\text{LogIC50})})} \]
Interpret the parameters

**IC50** is the concentration of agonist that gives a response half way between Bottom and Top. This is not the same as the response at Y=50. Depending on which units Y is expressed in, and the values of Bottom and Top, the IC50 may give a response nowhere near "50". Prism reports both the IC50 and its log.

**Top** and **Bottom** are plateaus in the units of the Y axis.

### 3.6.3.2 Equation: log(inhibitor) vs. response -- Variable slope

This equation is used when X values are logarithms of doses or concentrations. Use a related equation when X values are concentrations or doses.

**Introduction**

Many log(inhibitor) vs. response curves follow the familiar symmetrical sigmoidal shape. The goal is to determine the IC50 of the inhibitor - the concentration that provokes a response half way between the maximal (Top) response and the maximally inhibited (Bottom) response.

Many inhibitory dose-response curves have a standard slope of -1.0. This model does not assume a standard slope but rather fits the Hill Slope from the data, and so is called a **Variable slope** model. This is preferable when you have plenty of data points. It is also called a four-parameter dose-response curve, or four-parameter logistic curve, abbreviated 4PL.

**Step by step**

Create an XY data table. Enter the logarithm of the concentration of the inhibitor into X. Enter response into Y in any convenient units. Enter one data set into column A, and use columns B, C... for different treatments, if needed.

If you prefer to enter concentrations, rather than the logarithm of concentrations, use Prism to transform the X values to logs.
From the data table, click Analyze, choose nonlinear regression, choose the panel of equations "Dose-response curves - Inhibition" and then choose the equation "log(inhibitor) vs. response -- Variable slope".

If you have subtracted off any basal response, consider constraining Bottom to a constant value of 0.

**Model**

\[
Y = \text{Bottom} + \frac{\text{Top} - \text{Bottom}}{1 + 10^{((\text{LogIC50} - X) \times \text{HillSlope})}}
\]

**Interpret the parameters**

**IC50** is the concentration of agonist that gives a response half way between Bottom and Top. This is not the same as the response at \(Y=50\). Depending on which units \(Y\) is expressed in, and the values of Bottom and Top, the IC50 may give a response nowhere near "50". Prism reports both the IC50 and its log.

**HillSlope** describes the steepness of the family of curves. A HillSlope of -1.0 is standard, and you should consider constraining the Hill Slope to a constant value of -1.0. A Hill slope more negative than -1 (say -2) is steeper.

**Top** and **Bottom** are plateaus in the units of the Y axis.

**3.6.3.3 Equation: log(inhibitor) vs. normalized response**

This equation is used when X values are logarithms of doses or
concentrations. Use a related equation when X values are concentrations or doses.

Introduction

Many log(inhibitor) vs. response curves follow the familiar symmetrical sigmoidal shape.

If you have good control data, it can make sense to normalize the response to run between 0% and 100%. This model assumes that the data have been normalized, so forces the curve to run from 100% down to 0%. The goal is to determine the IC50 of the inhibitor - the concentration that provokes a response equal to 50%.

It only makes sense to fit a normalized model when you are sure you have defined 0% and 100% quite accurately. If your data define a complete sigmoidal curve, it is best to fit the entire curve and let Prism fit the Top and Bottom plateaus. If your data don't form a full sigmoidal curve, but you can define the bottom and top by solid control data, then fitting to a normalized model is preferable.

This model assumes that the dose response curve has a standard slope, equal to a Hill slope (or slope factor) of -1.0. This is the slope expected when a ligand binds to a receptor following the law of mass action, and is the slope expected of a dose-response curve when the second messenger created by receptor stimulation binds to its receptor by the law of mass action. If you don't have many data points, consider using the standard slope model. If you have lots of data points, pick the variable slope model to determine the Hill slope from the data.

Step by step

Create an XY data table. Enter the logarithm of the concentration of the inhibitor into X. Enter response into Y in any convenient units. Enter one data set into column A, and use columns B, C... for different treatments, if needed.

If you prefer to enter concentrations, rather than the logarithm of concentrations, use Prism to transform the X values to logs.

From the data table, click Analyze, choose nonlinear regression, choose the panel of equations "Dose-response curves - Inhibition" and then choose the equation "log(inhibitor) vs. normalized response".
Model

\[ Y = \frac{100}{1 + 10^{(X - \text{LogIC50})}} \]

Interpret the parameter

**IC50** is the concentration of agonist that gives a response half way between Bottom and Top. This is not the same as the response at \( Y = 50 \). Depending on which units \( Y \) is expressed in, and the values of Bottom and Top, the IC50 may give a response nowhere near "50". Prism reports both the IC50 and its log.

3.6.3.4 Equation: log(inhibitor) vs. normalized response -- Variable slope

This equation is used when \( X \) values are logarithms of doses or concentrations. **Use a related equation** \[317\] when \( X \) values are concentrations or doses.

Introduction

Many log(inhibitor) vs. response curves follow the familiar symmetrical sigmoidal shape.

If you have good control data, it can make sense to **normalize** the response to run between 0% and 100%. This model assumes that the data have been normalized, so forces the curve to run from 100% down to 0%. The goal is to determine the IC50 of the inhibitor - the concentration that provokes a response equal to 50%.
It only makes sense to fit a normalized model when you are sure you have defined 0% and 100% quite accurately. If your data define a complete sigmoidal curve, it is best to fit the entire curve and let Prism fit the Top and Bottom plateaus. If your data don't form a full sigmoidal curve, but you can define the bottom and top by solid control data, then fitting to a normalized model is preferable.

Many inhibitory dose-response curves have a standard slope of -1.0. This model does not assume a standard slope but rather fits the Hill Slope from the data, and so is called a Variable slope model. This is preferable when you have plenty of data points.

**Step by step**

Create an XY data table. Enter the logarithm of the concentration of the inhibitor into X. Enter response into Y in any convenient units. Enter one data set into column A, and use columns B, C... for different treatments, if needed.

If you prefer to enter concentrations, rather than the logarithm of concentrations, use Prism to transform the X values to logs.

From the data table, click Analyze, choose nonlinear regression, choose the panel of equations "Dose-response curves - Inhibition" and then choose the equation "log(inhibitor) vs. normalized response -- variable slope".

**Model**

\[ Y = \frac{100}{1 + 10^{((\log IC50 - X) \times HillSlope)}} \]
Interpret the parameters

**IC50** is the concentration of agonist that gives a response half way between Bottom and Top. This is not the same as the response at Y=50. Depending on which units Y is expressed in, and the values of Bottom and Top, the IC50 may give a response nowhere near "50". Prism reports both the IC50 and its log.

**HillSlope** describes the steepness of the family of curves. A HillSlope of -1.0 is standard, and you should consider constraining the Hill Slope to a constant value of -1.0. A Hill slope more negative than -1 (say -2) is steeper.

3.6.3.5 Equation: [Inhibitor] vs. response

This equation is used when X values are concentrations or doses. Use a related equation when X values are logarithms of concentrations or doses.

Introduction

Many log(inhibitor) vs. response curves follow the familiar symmetrical sigmoidal shape. The goal is to determine the IC50 of the inhibitor - the concentration that provokes a response half way between the maximal (Top) response and the maximally inhibited (Bottom) response.

This model assumes that the dose response curves has a **standard slope**, equal to a Hill slope (or slope factor) of -1.0. This is the slope expected when a ligand binds to a receptor following the law of mass action, and is the slope expected of a dose-response curve when the second messenger created by receptor stimulation binds to its receptor by the law of mass action. If you don't have many data points, consider using the standard slope model. If you have lots of data points, pick the variable slope model to determine the Hill slope from the data.

Step by step

Create an XY data table. Enter the concentrations of the inhibitor into X. Enter response into Y in any convenient units. Enter one data set into column A, and use columns B, C... for different treatments, if needed.
From the data table, click Analyze, choose nonlinear regression, choose the panel of equations "Dose-response curves - Inhibition" and then choose the equation "[Inhibitor] vs. response".

If you have subtracted off any basal response, consider constraining Bottom to a constant value of 0.

Since the uncertainty of the EC50 is very asymmetric, be sure to choose to compute the confidence intervals using the likelihood ratio asymmetric method.

Double click on the X axis of the graph, and choose (at the upper left of the Format Graph dialog) to stretch the axis to a logarithmic scale.

Model

\[ Y = \text{Bottom} + \frac{(\text{Top} - \text{Bottom})}{(1 + \frac{X}{\text{IC50}})} \]

Interpret the parameters

\textbf{IC50} is the concentration of agonist that gives a response half way between Bottom and Top. This is not the same as the response at \(Y=50\). Depending on which units \(Y\) is expressed in, and the values of Bottom and Top, the IC50 may give a response nowhere near "50". Prism reports both the IC50 and its log.

\textbf{Top} and \textbf{Bottom} are plateaus in the units of the Y axis.
3.6.3.6 Equation: [Inhibitor] vs. response -- Variable slope

This equation is used when X values are concentrations or doses. Use a related equation when X values are logarithms of concentrations or doses.

Introduction

Many log(inhibitor) vs. response curves follow the familiar symmetrical sigmoidal shape. The goal is to determine the IC50 of the inhibitor - the concentration that provokes a response half way between the maximal (Top) response and the maximally inhibited (Bottom) response.

Many inhibitory dose-response curves have a standard slope of -1.0. This model does not assume a standard slope but rather fits the Hill Slope from the data, and so is called a Variable slope model. This is preferable when you have plenty of data points. It is also called a four-parameter dose-response curve, or four-parameter logistic curve, abbreviated 4PL.

Step by step

Create an XY data table. Enter the concentrations of the inhibitor into X. Enter response into Y in any convenient units. Enter one data set into column A, and use columns B, C... for different treatments, if needed.

From the data table, click Analyze, choose nonlinear regression, choose the panel of equations "Dose-response curves - Inhibition" and then choose the equation "[Inhibitor] vs. response -- Variable slope".

If you have subtracted off any basal response, consider constraining Bottom to a constant value of 0.

Since the uncertainty of the EC50 is very asymmetric, be sure to choose to compute the confidence intervals using the likelihood ratio asymmetric method.

Double click on the X axis of the graph, and choose (at the upper left of the Format Graph dialog) to stretch the axis to a logarithmic scale.

Model

\[ Y = \text{Bottom} + \frac{(\text{Top} - \text{Bottom})}{(1 + \left(\frac{\text{X}^{\text{HillSlope}}}{\text{IC50}^{\text{HillSlope}}}\right))} \]
Interpret the parameters

**IC50** is the concentration of agonist that gives a response half way between Bottom and Top. This is not the same as the response at Y=50. Depending on which units Y is expressed in, and the values of Bottom and Top, the IC50 may give a response nowhere near "50". Prism reports both the IC50 and its log.

**HillSlope** describes the steepness of the family of curves. A HillSlope of -1.0 is standard, and you should consider constraining the Hill Slope to a constant value of -1.0. A Hill slope more negative than -1 (say -2) is steeper.

**Top** and **Bottom** are plateaus in the units of the Y axis.

3.6.3.7 Equation: [Inhibitor] vs. normalized response

This equation is used when X values are concentrations or doses. Use a related equation when X values are logarithms of concentrations or doses.

**Introduction**

Many log(inhibitor) vs. response curves follow the familiar symmetrical sigmoidal shape.

If you have good control data, it can make sense to **normalize** the response to run between 0% and 100%. This model assumes that the
data have been normalized, so forces the curve to run from 100% down to 0%. The goal is to determine the IC50 of the inhibitor - the concentration that provokes a response equal to 50%.

It only makes sense to fit a normalized model when you are sure you have defined 0% and 100% quite accurately. If your data define a complete sigmoidal curve, it is best to fit the entire curve and let Prism fit the Top and Bottom plateaus. If your data don't form a full sigmoidal curve, but you can define the bottom and top by solid control data, then fitting to a normalized model is preferable.

This model assumes that the dose response curve has a standard slope, equal to a Hill slope (or slope factor) of -1.0. This is the slope expected when a ligand binds to a receptor following the law of mass action, and is the slope expected of a dose-response curve when the second messenger created by receptor stimulation binds to its receptor by the law of mass action. If you don't have many data points, consider using the standard slope model. If you have lots of data points, pick the variable slope model to determine the Hill slope from the data.

**Step by step**

Create an XY data table. Enter the concentrations of the inhibitor into X. Enter response into Y in any convenient units. Enter one data set into column A, and use columns B, C... for different treatments, if needed.

From the data table, click Analyze, choose nonlinear regression, choose the panel of equations "Dose-response curves - Inhibition" and then choose the equation "[Inhibitor] vs. normalized response".

Since the uncertainty of the EC50 is very asymmetric, be sure to choose to compute the confidence intervals using the likelihood ratio asymmetric method.

Double click on the X axis of the graph, and choose (at the upper left of the Format Graph dialog) to stretch the axis to a logarithmic scale.

**Model**

\[ Y = \frac{100}{1+X/IC50} \]
Interpret the parameter

**IC50** is the concentration of agonist that gives a response half way between Bottom and Top. This is not the same as the response at Y=50. Depending on which units Y is expressed in, and the values of Bottom and Top, the IC50 may give a response nowhere near "50". Prism reports both the IC50 and its log.

3.6.3.8 Equation: [Inhibitor] vs. normalized response -- Variable slope

This equation is used when X values are concentrations or doses. Use a related equation when X values are logarithms of concentrations or doses.

Introduction

Many log(inhibitor) vs. response curves follow the familiar symmetrical sigmoidal shape.

If you have good control data, it can make sense to normalize the response to run between 0% and 100%. This model assumes that the data have been normalized, so forces the curve to run from 100% down to 0%. The goal is to determine the IC50 of the inhibitor - the concentration that provokes a response equal to 50%.

It only makes sense to fit a normalized model when you are sure you have defined 0% and 100% quite accurately. If your data define a complete sigmoidal curve, it is best to fit the entire curve and let Prism
fit the **Top and Bottom plateaus** \[^{[60]}\]. If your data don't form a full sigmoidal curve, but you can define the bottom and top by solid control data, then fitting to a normalized model is preferable.

Many inhibitory dose-response curves have a standard slope of -1.0. This model does not assume a standard slope but rather fits the Hill Slope from the data, and so is called a **Variable slope** model. This is preferable when you have plenty of data points.

**Step by step**

Create an XY data table. Enter the concentrations of the inhibitor into X. Enter response into Y in any convenient units. Enter one data set into column A, and use columns B, C... for different treatments, if needed.

From the data table, click Analyze, choose nonlinear regression, choose the panel of equations "Dose-response curves - Inhibition" and then choose the equation "\( \log(\text{inhibitor}) \) vs. normalized response -- variable slope".

Since the uncertainty of the EC50 is very asymmetric, be sure to choose to compute the confidence intervals using the **likelihood ratio asymmetric method** \[^{[88]}\].

Double click on the X axis of the graph, and choose (at the upper left of the Format Graph dialog) to stretch the axis to a logarithmic scale.

**Model**

\[ Y = \frac{100}{1 + \left(\frac{X^{\text{HillSlope}}}{\text{IC50}^{\text{HillSlope}}} \right)} \]
Interpret the parameters

**IC50** is the concentration of agonist that gives a response half way between Bottom and Top. This is not the same as the response at Y=50. Depending on which units Y is expressed in, and the values of Bottom and Top, the IC50 may give a response nowhere near "50". Prism reports both the IC50 and its log.

**HillSlope** describes the steepness of the family of curves. A HillSlope of -1.0 is standard, and you should consider constraining the Hill Slope to a constant value of -1.0. A Hill slope more negative than -1 (say -2) is steeper.

3.6.4 Dose-response -- Special

3.6.4.1 Asymmetrical (five parameter)

Introduction

The standard dose-response curve is sometimes called the **four-parameter logistic equation**. It fits the bottom and top plateaus of the curve, the EC50, and the slope factor (Hill slope). This curve is symmetrical around its midpoint. To extend the model to handle curves that are not symmetrical, the Richards equation adds an additional parameter, S, which quantifies the asymmetry. This equation is sometimes referred to as a **five-parameter logistic equation**, abbreviated 5PL.

Step by step

Create an XY data table. Enter the logarithm of the concentration of the agonist into X. Enter response into Y in any convenient units.

From the data table, click Analyze, choose nonlinear regression, and choose the panel of equations: Dose-Response -- Special. Then choose **Asymmetrical (five parameter)**.

Consider constraining the Hill Slope to a constant value of 1.0 (stimulation) or -1 (inhibition).
Also consider whether Bottom or Top should be fixed to constant values, or shared between data sets.

**Model**

\[
\text{LogXb} = \text{LogEC50} + \frac{1}{\text{HillSlope}} \times \log\left(\frac{2^{\frac{1}{S}} - 1}{1 + 10^{\left(\text{LogXb} - X\right) \times \text{HillSlope}}}\right)
\]

Numerator = Top - Bottom

Denominator = \left(1 + 10^{\left(\text{LogXb} - X\right) \times \text{HillSlope}}\right)^S

\[
Y = \text{Bottom} + \frac{\text{Numerator}}{\text{Denominator}}
\]

**Interpret the parameters**

**Bottom** and **Top** are the plateaus at the left and right ends of the curve, in the same units as Y.

**LogEC50** is the concentrations that give half-maximal effects, in the same units as X. Note that the logEC50 is not the same as the inflection point Xb (see below).

**HillSlope** is the unitless slope factor or Hill slope. Consider constraining it to equal 1.0 (stimulation) or -1 (inhibition).

**S** is the unitless symmetry parameter. If S=1, the curve is symmetrical and identical to the standard dose-response equation. If S is distinct than 1.0, then the curve is asymmetric as shown below.
3.6.4.2 Equation: Biphasic dose-response

Introduction

A common deviation from the standard monotonic sigmoid shape is the biphasic sigmoid shape.

Step by step

Create an XY data table. Enter the logarithm of the concentration of the agonist into X. Enter response into Y in any convenient units.

From the data table, click Analyze, choose nonlinear regression, and choose the panel of equations: Dose-Response -- Special. Then choose Biphasic dose-response.
Consider constraining \( nH1 \) and \( nH2 \) to constant values of 1.0 (stimulation) or -1 (inhibition).

Also consider whether Bottom or Top should be fixed to constant values, or shared between data sets.

**Model**

\[
\text{Span} = \text{Top} - \text{Bottom} \\
\text{Section1} = \text{Span} \times \text{Frac} / (1 + 10^{((\text{LogEC50}_1 - X) \times nH1)}) \\
\text{Section2} = \text{Span} \times (1 - \text{Frac}) / (1 + 10^{((\text{LogEC50}_2 - X) \times nH2)}) \\
Y = \text{Bottom} + \text{Section1} + \text{Section2}
\]

**Interpret the parameters**

*Bottom* and *Top* are the plateaus at the left and right ends of the curve, in the same units as \( Y \).

*LogEC50_1* and *LogEC50_2* are the concentrations that give half-maximal stimulatory and inhibitory effects in the same units as \( X \).

*nH1* and *nH2* are the unitless slope factors or Hill slopes. Consider constraining these to equal 1.0 (stimulation) and -1 (inhibition).

*Frac* is the proportion of maximal response due to the more potent phase.
3.6.4.3 Equation: Bell-shaped dose-response

Introduction

Some drugs may cause an inhibitory response at low concentrations, and a stimulatory response at high concentrations, or vice-versa. The net result is a bell-shaped dose-response curve.

The model explained here is the sum of two dose-response curves, one that stimulates and one that inhibits. you will need lots of data to determine all the parameters without ambiguity, so this model will rarely be useful for data analysis. But it might be useful as a way to draw a smooth curve through the data

Step by step

Create an XY data table. Enter the logarithm of the concentration of the agonist into X. Enter response into Y in any convenient units.

From the data table, click Analyze, choose nonlinear regression, and choose the panel of equations: Dose-Response -- Special. Then choose Bell-shaped dose-response.

Consider constraining nH1 and nH2 to constant values of 1.0 (stimulation) and -1 (inhibition).

Model

\[
\begin{align*}
\text{Span}_1 &= \text{Plateau}_1 - \text{Dip} \\
\text{Span}_2 &= \text{Plateau}_2 - \text{Dip} \\
\text{Section}_1 &= \frac{\text{Span}_1}{(1+10^{((\text{LogEC50}_1 - X) \times nH1)})} \\
\text{Section}_2 &= \frac{\text{Span}_2}{(1+10^{((X - \text{LogEC50}_2) \times nH2)})} \\
Y &= \text{Dip} + \text{Section}_1 + \text{Section}_2
\end{align*}
\]
Interpret the parameters

**Plateau1** and **Plateau2** are the plateaus at the left and right ends of the curve, in the same units as Y.

**Dip** is the plateau level in the middle of the curve, in same units as Y. Note that when the curve goes up first (then down), this parameter named *Dip* would better be named *Peak*.

**LogEC50_1** and **LogEC50_2** are the concentrations that give half-maximal stimulatory and inhibitory effects in the same units as X.

**nH1** and **nH2** are the unitless slope factors or Hill slopes. Consider constraining these to equal 1.0 (stimulation) and -1 (inhibition).

3.6.4.4 Equation: Operational model - Depletion

**Introduction to the operational model**

The EC50 is determined by two properties of the agonist:

- How well it binds to the receptor, quantified by the **affinity** of the drug for binding to its receptor.
- How well it causes a response once bound. This property is known as the agonist’s **efficacy**. Since efficacy depends on both agonist and tissue, a single drug acting on a single kind of receptor can have different efficacies, and thus different EC50 values, in different tissues.

A single dose-response experiment cannot determine affinity and efficacy. A drug that binds with high affinity but has low efficacy will produce exactly the same dose-response curve as a drug with low affinity and high efficacy.

To untangle affinity from efficacy, globally fit a dose-response curve of a full agonist and a second dose-response curve determined after treating the cells or tissue with an alkylating agent (or some other irreversible treatment) that reduces the number of accessible receptors. With fewer receptors, the dose response curve is shifted down and usually to the right.

The operational model assumes that the affinity of the drug for the receptors is not altered by reducing the number of available receptors. It also assumes that the maximum possible response in the tissue remains unchanged (the treatment was specific for the receptors you are studying). Accepting these assumptions, fitting the operational model globally will determine the affinity of the agonist for the receptors.

**Step by step**

Create an XY data table. Enter the logarithm of the concentration of the agonist ligand into X. Enter response into Y in any convenient units. Enter data with a full agonist and no receptor depletion into column A. Enter data collected after receptor depletion into column B. Repeat, if you have data with different levels of receptor depletion for column C, D, E, ... You don't have to know the degree to which the receptors are depleted, and don't have to enter any values in the column titles (although they are useful as labels).

From the data table, click Analyze, choose nonlinear regression, and choose the panel of equations: Dose-Response -- Special. Then choose *Operational Model - Depletion*.

If you have subtracted off any basal response, consider constraining the parameter Basal to a constant value of zero.
Also consider constraining the transducer slope n to a constant value of 1.0. When set to 1.0, all dose-response curves are constrained to have Hill slopes of 1.0, which is observed commonly.

Model

\[
\text{operate} = \frac{\left(10^{\log KA} + 10^X\right)}{10^{\log \tau + X}} \quad n
\]

\[
Y = \text{Basal} + \frac{\text{Effectmax} - \text{Basal}}{1 + \text{operate}}
\]

Interpret the parameters

**Effectmax** is the maximum possible system response, in units of the Y axis. It is the top plateau of the dose-response curve obtained with a full agonist without receptor depletion. If your agonist isn't a full agonist, the EffectMax might be higher than the top plateau of the no depletion curve.

**Basal** is the response in absence of agonist, in same units as Y. If you have subtracted off any basal response, constrain basal to a constant value of zero.

**KA** is the agonist-receptor dissociation constant, in same units as X (usually molar). It measures the affinity of the full agonist for the receptors, which is the main goal of this kind of experiment. Prism reports both KA and its logarithm. It is not the same as the EC50.

**tau** is the transducer constant, a practical measure of efficacy. It is the inverse of the fraction of receptors that must be occupied by agonist to obtain the half-maximal response. If tau equals 10, that means that occupation of only 10% of the receptors leads to a half-maximal
response. If tau equals 1.0, that means that it requires occupation of all the receptors to give a half-maximal response. This would happen in a tissue where the receptors had been substantially depleted. Because \( t \) is a property of both the tissue and receptor system, it is not a direct measure of intrinsic efficacy, which is commonly defined as a property belonging only to an agonist-receptor pair, irrespective of the assay system in which it is measured. Prism reports both tau and its logarithm for each data set.

\( n \) is the Unitless transducer slope. It is similar to, but not identical to, the Hill slope. In most cases, \( n \) is constrained to a constant value of 1.0, in which case all the dose-response curves will have Hill slopes of 1.0. If \( n \) does not equal 1.0, the Hill Slope does not equal either 1.0 or \( n \).

**Notes**

Since Tau measures efficacy, Prism fits a different value of tau for each data set. Receptor depletion reduce the value of tau. The other parameters are fit globally, to find one best-fit value for all the data sets.

<table>
<thead>
<tr>
<th>Reference</th>
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**3.6.4.5 Equation: Operational model - Partial agonist**

**Introduction to the operational model**

The EC50, fit by standard dose-response models, is determined by two properties of the agonist:

- How well it binds to the receptor, quantified by the **affinity** of the drug for binding to its receptor.

- How well it causes a response once bound. This property is known as the agonist’s **efficacy**. Since efficacy depends on both agonist and tissue, a single drug acting on a single kind of receptor can have different efficacies, and thus different EC50 values, in different tissues.

A single dose-response experiment cannot determine affinity and efficacy. A drug that binds tightly with high affinity but has low efficacy,
will produce exactly the same dose-response curve as a drug with low affinity and high efficacy.

To determine the affinity of a partial agonist, use the operational model to globally fit the dose-response curves of both a full agonist and the partial agonist. The data from the full-agonist determines the maximum possible effect. Knowing that, the fitting can determine the affinity of the partial agonist.

**Step by step**

Create an XY data table. Enter the logarithm of the concentration of the agonist ligand into X. Enter response into Y in any convenient units. Enter data with the full agonist into column A. Enter data collected with a partial agonist into column B. Repeat, if you have data with different partial agonists, for column C, D, E, ..., each with a different amount of depletion.

From the data table, click Analyze, choose nonlinear regression, and choose the panel of equations: Dose-Response -- Special. Then choose *Operational Model - Partial agonists*.

If you have subtracted off any basal response, consider constraining the parameter Basal to a constant value of zero.

Also consider constraining the transducer slope n to a constant value of 1.0. When set to 1.0, all dose-response curves are constrained to have Hill slopes of 1.0, which is observed commonly. If n is not 1.0, the Hill slopes will not be 1.0, but the Hill slopes will not equal exactly n.

**Model**

\[
\text{operate} = \frac{10^{\text{log} K_A} + (10^X)}{10^{\text{log} \tau + X}}^n
\]

\[
\text{<A> } Y = \text{Basal} + \frac{\text{Effectmax-Basal}}{1 + 10^{((\text{LogEC50}-X)*n)}}
\]

\[
\text{<~A> } Y = \text{Basal} + \frac{\text{Effectmax-Basal}}{1 + \text{operate}}
\]

The second line is preceded with <A> which means it only applies to the first data set. It fits a variable slope dose-response curve. The third line is preceded with <~A> which means it applies to all data set except the first. It fits the operational model to determine the affinity (KA) of the partial agonist.
Interpret the parameters

**Effectmax** is the maximum possible system response, in units of the Y axis. It is the top plateau of the full agonist's dose-response curve.

**Basal** is the response in absence of agonist, in same units as Y. If you have subtracted off any basal response, constrain basal to a constant value of zero.

**KA** is the equilibrium dissociation constant of the partial agonist(s), in same units as X (usually molar). It measures the affinity of the partial agonist for the receptors, which is the main goal of this kind of experiment. Prism reports both KA and its logarithm. It is not the same as the EC50.

**tau** is the transducer constant, a practical measure of efficacy. It is the inverse of the fraction of receptors that must be occupied by agonist to obtain the half-maximal response. If t equals 10, that means that occupation of only 10% of the receptors leads to a half-maximal response. If t equals 1.0, that means that it requires occupation of all the receptors to give a half-maximal response. This would happen with a partial agonist. Prism reports both tau and its logarithm, and fits tau individually for each data set.

**n** is the Unitless transducer slope. It is similar to, but not identical to, the Hill slope. In most cases, n is constrained to a constant value of 1.0, in which case all the dose-response curves will have Hill slopes of 1.0. If n does not equal 1.0, the Hill Slope does not equal either 1.0 or n.
3.6.4.6 Equation: Gaddum/Schild EC50 shift

**Introduction**

A competitive inhibitor competes for agonist binding to a receptor, and shifts the dose-response curve to the right without changing the maximum response. By fitting all the curves globally, you can determine the affinity of the competitive inhibitor.

**Step by step**

1. Create an XY data table with subcolumns to match your experimental design.

2. Enter the logarithm of the concentration of the agonist ligand into X. For example, if a concentration is 1nM, enter -9. Don't enter 1e-9.

3. Enter response into Y in any convenient units. Enter data with no inhibitor into column A. Enter data collected with a constant concentration of inhibitor into column B. Repeat, if you have data, for column C, D, E, ..., each with a different concentration of inhibitor.

4. Enter the inhibitor concentration in molar as the column titles of the data table. If the concentration is 1nM enter '1e-9' or '0.00000001' into the column titles. Don't enter '-9'. Don't forget to enter '0' as the column title for data set A, since these are the control data with no inhibitor. These values entered into the column title are used in the analysis; they are more than just labels, so need to be entered in the correct format.

5. From the data table, click Analyze, choose nonlinear regression, and choose the panel of equations: Dose-Response -- Special. Then choose Gaddum/Schild EC50 shift.

6. Consider constraining the parameters HillSlope and SchildSlope to their standard values of 1.0. This is especially useful if you don't have many data points, and therefore cannot fit these parameters.

Reference

Notes on units

- This equation was written so the X values are entered as logarithm of concentration of the agonist, while the column titles are entered as concentration of antagonist. This is simply the way this equation was written. It would be easy enough to clone the equation and write it with different conventions if you wanted to.

- The EC50 is reported in the antilog of the units you used to enter X values. If you entered X as log(molar), then the EC50 will be in molar. If you entered X values as log(dose in milligrams), then the EC50 will be in milligrams.

- The pA2 is the negative logarithm of the concentration of antagonist needed to shift the curve by a factor of 2. That concentration is in the same units you used to enter the antagonist concentration into the column titles. If you entered concentration in nM into the column titles, the pA2 is the negative logarithm of nM. If you entered concentration as a dose in mg/kg, then the pA2 is the negative logarithm of mg/kg.

- If you want to transform the pA2 to a concentration, first flip the sign (multiply by -1) and then take the antilog (ten to that power). If you also do so for the confidence limits, note that flipping the sign also changes the order of the limits, so the "lower" confidence limit is actually the upper limit, and the "upper" limit is actually the lower limit.

Model

\[
\begin{align*}
EC50 &= 10^{\text{LogEC50}} \\
Antag &= 1 + \frac{B}{(10^{-1 \times \text{pA2}})}^{\text{SchildSlope}} \\
\text{LogEC} &= \text{Log}(EC50 \times \text{Antag}) \\
Y &= \text{Bottom} + \frac{\text{Top} - \text{Bottom}}{1 + 10^{((\text{LogEC} - X) \times \text{HillSlope})}}
\end{align*}
\]
Interpret the parameters

**EC50** is the concentration of agonist that gives half maximal response in the absence of inhibitor. Prism reports both the EC50 and its log. The logEC50 is in the same units you used to enter the X values. The EC50 is in the antilog of those units.

**pA2** is the negative logarithm of the concentration of antagonist needed to shift the dose response curve by a factor of 2. If the HillSlope and SchildSlope are fixed to 1.0, it is the pKb, the negative log of the equilibrium dissociation constant (Molar) of inhibitors binding to the receptors. The units are the negative log of whatever units you used to enter concentrations as column titles on the data table.

**HillSlope** describes the steepness of the family of curves. A HillSlope of 1.0 is standard, and you should consider constraining the Hill Slope to a constant value of 1.0.

**SchildSlope** quantifies how well the shifts correspond to the prediction of competitive interaction. If the competitor is competitive, the SchildSlope will equal 1.0. You should consider constraining SchildSlope to a constant value of 1.0. antagonist term, [B], is now raised to the power S, where S denotes the Schild slope factor. If the shift to the right is greater than predicted by competitive interactions, S will be greater
than 1. If the rightward shift is less than predicted by competitive interaction, then S will be less than 1.

**Top** and **Bottom** are plateaus in the units of the Y axis.

Notes:

- All six of these are shared among all the data sets, so you will only see one best fit value for each parameter, not one for each data set.

- Prism does not report the EC50 for each curve, only the EC50 for the first curve (the one without any antagonist).

- The variable B in the model is defined to be a data set constant whose value comes from the column titles. The results page will show these values of B. Make sure they are the concentration of antagonist used in each data set (and not the log of concentrations).

**When is the Schild model valid?**

Colquhoun (1) has shown that the Schild model is valid whenever you can make these assumptions:

- The antagonist, B, is a true antagonist that, alone, does not change the conformation of the receptor.

- Binding of agonist, A, and antagonist, B, is mutually exclusive at every binding site.

- B has the same affinity for every binding site.

- The observed response is the same if the occupancy of each site by A is the same, regardless of how many sites are occupied by B.

- Measurements are made at equilibrium.

3.6.4.7 Equation: EC50 shift

Introduction

An competitive inhibitor competes for agonist binding to a receptor, and shifts the dose-response curve to the right without changing the maximum response. This model fits the two dose response curves and determines the fold shift.

Step by step

1. Create an XY data table.

2. Enter the logarithm of the concentration of the agonist ligand into the X column.

3. Enter response into Y in any convenient units. Enter data with no inhibitor into column A. Enter data collected with a constant concentration of inhibitor into column B. You can label columns A and B with any column title you like. The column title does not get used as part of the analysis.

4. From the data table, click Analyze, choose nonlinear regression, and choose the panel of equations: Dose-Response -- Special. Then choose Dose shift.

5. If you have subtracted off any baseline signal, constrain the parameter Bottom to a constant value of zero.

Model

\[ \text{LogEC} = \text{LogEC50Control} \]
\[ \text{LogEC} = \text{LogEC50Control} + \log(\text{EC50Ratio}) \]
\[ Y = \text{Bottom} + \left( \text{Top} - \text{Bottom} \right) / \left( 1 + 10^{(\text{LogEC} - X) \times \text{HillSlope}} \right) \]
**EC50Control** is the concentration of agonist that gives half maximal response in the absence of modulator.

**Top** and **Bottom** are plateaus in the units of the Y axis (shared).

**EC50Ratio** is the ratio of EC50 in presence of inhibitor divided by EC50 of agonist alone.

**HillSlope** is the slope factor (shared)

**Notes**

- If you have several concentrations of antagonist, use a different model that will directly fit the Schild model and determine the pA2.

- If you want to test the null hypothesis that the EC50 ratio is 1.0 (no shift), that is easy to do. Go to the Compare tab, choose to test whether a selected parameter differs from a hypothetical value, choose the parameter EC50Ratio, and enter the hypothetical value 1.0.

- This model shares Top, Bottom and HillSlope. So it assumes that both models are exactly the same except (perhaps) for the EC50. You can go to the Constrain tab and lift this constraint for any or all of these three parameters, if you think it makes sense not to share them.
### 3.6.4.8 Equation: Allosteric EC50 shift

#### Introduction

An allosteric modulator can reduce or enhance agonist binding. This model fits entire dose-response curves determined in the absence and presence of a modulator. The goal is to learn the affinity of the modulator for binding to its site, and also determine the value of alpha, the ternary complex constant that quantifies the degree to which binding of the modulator alters the affinity of the radioligand for the receptor site.

#### Step by step

1. Create an XY data table.

2. Enter the logarithm of the concentration of the agonist ligand into X. If the concentration is 1nM, enter -9.

3. Enter response into Y in any convenient units. Enter data with no modulator into column A. Enter data collected with a constant concentration of modulator into column B. Repeat, if you have data, for column C, D, E, ..., each with a different concentration of modulator.

4. Enter the modulator concentration (in molar so 1nM is entered as '1e-9') into the column titles. Don't forget to enter '0' as the column title for data set A. These column titles are not just labels. The values you enter become part of the analysis.

5. From the data table, click Analyze, choose nonlinear regression, and choose the panel of equations: Dose-Response -- Special. Then choose *Allosteric EC50 shift*.

You do not need to constrain any parameters to constant values

#### Model

```
EC50=10^LogEC50
KB=10^LogKB
alpha=10^Logalpha
Antag=(1+B/KB)/(1+alpha*B/KB)
LogEC=Log(EC50*Antag)
Y=Bottom+(Top-Bottom)/(1+10^((LogEC-X)*HillSlope))
```
EC50 is the concentration of agonist that gives half maximal response in the absence of modulator.

Kb is the equilibrium dissociation constant (Molar) of modulator binding to its allosteric site. It is in the same molar units used to enter the modulator concentration into column titles on the data table.

Alpha is the ternary complex constant. When alpha=1.0, the modulator won't alter binding. If alpha is less than 1.0, then the modulator reduces ligand binding. If alpha is greater than 1.0, then the modulator increases binding. In the example shown about, alpha equals 0.01 so the modulator greatly decreases binding.

Top and Bottom are plateaus in the units of the Y axis.

Notes

- This model is designed to analyze data when the modulator works via an allosteric site. Since the agonist and modulator are acting via different sites, it is incorrect to refer to the modulator as a competitor.

- The model is written to fit the logarithm of alpha, rather than alpha itself. This is because alpha is asymmetrical: All values from 0 to 1 mean that the modulator decreases binding, while all values from 1 to infinity mean that the modulator enhances binding. On a log scale, its values are more symmetrical, so the confidence interval computed on
a log scale (as Prism does) are more accurate. Prism reports both alpha and log(alpha).

- This model assumes that the allosteric modulator is present in excess, so the concentration you added is very close to its free concentration. This model won’t work when the concentration of allosteric modulator is limiting (as it is when G proteins alter agonist binding to many receptors). No explicit model can handle this situation. You need to define the model with an implicit equation (Y on both sides of the equals sign) and Prism cannot handle such equations.

- Note two important points about the progression of curves from left to right with increasing concentrations of the allosteric modulator. First, note that the maximum response doesn’t change. Second, note that the effect of the modulator to right-shift the dose-response curve reaches a maximum as the modulator saturates its binding site.

Reference


3.6.4.9 Equation: ECanything

Introduction

Many log(dose) response curves follow the familiar symmetrical sigmoidal shape. The usual goal is to determine the EC50 of the agonist - the concentration that provokes a response half way between the basal (Bottom) response and the maximal (Top) response. But you can determine any spot along the curve, say a EC80 or EC90.

Many dose-response curves have a standard slope of 1.0. This model does not assume a standard slope but rather fits the Hill Slope from the data. Hence the name **Variable slope** model. This is preferable when you have plenty of data points.

Step by step

Create an XY data table. Enter the logarithm of the concentration of the agonist into X. Enter response into Y in any convenient units. Enter one data set into column A, and use columns B, C... for different treatments, if needed.
From the data table, click Analyze, choose nonlinear regression, and choose the panel of equations: Dose-Response -- Special. Then choose "log(Agonist) vs. response -- Find ECanything".

You must constrain the parameter F to have a constant value between 0 and 100. Set F to 80 if you want to fit the EC80. If you constrain F to equal 50, then this equation is the same as a variable slope dose-response curve.

Consider constraining the parameter HillSlope to its standard value of 1.0 or -1. This is especially useful if you don't have many data points, and therefore cannot fit the slope very well.

If you have subtracted off any basal response, consider constraining Bottom to a constant value of 0.

**Model**

\[
\log(\text{EC50}) = \log(\text{ECF}) - \frac{1}{\text{HillSlope}} \log\left(\frac{F}{100-F}\right)
\]

\[
Y = \text{Bottom} + \frac{\text{Top} - \text{Bottom}}{1 + 10^{\left(\log(\text{EC50}) - X\right) \cdot \text{HillSlope}}}
\]

**Interpret the parameters**

**ECF** is the concentration of agonist that gives a response F percent of the way between Bottom and Top. Prism reports both the ECF and its log.

**HillSlope** describes the steepness of the family of curves. A HillSlope of 1.0 is standard, and you should consider constraining the Hill Slope to a constant value of 1.0. A Hill slope greater than 1.0 is steeper, and a Hill slope less than 1.0 is shallower.

**Top** and **Bottom** are plateaus in the units of the Y axis.
Adapting this equation

ICAnything

This equation can also fit inhibitory data where the curve goes downhill rather than uphill. The best-fit value of the Hill Slope will be negative in this case. The result will always be reported as ECF. Let's say you set F=80. Then the ECF for inhibitory data would be the concentration (X value) required to bring the curve down to 80%. If you want the concentration that brings the curve down by 80%, to 20%, then you'd need to set F equal to 20. If you want the result to say "ICF" rather than "ECF", clone the equation and then edit to make that change.

X is concentration rather than log concentration

The dose response equations where X is dose or concentration (rather than its logarithm) can be easily adapted to fit the ECF or ICF. Simply clone the equation and add one line to the top that turns EC50 into an intermediate variable and adds the parameters ECF (which you'll fit) and F (which you'll set as a constant). For example the [Agonist] vs. response -- variable slope equation becomes:

\[
EC50 = ECF/((F/(100-F))^{(1/HillSlope)})
\]

\[
Y=Bottom + (X^{Hillslope})*(Top-Bottom)/(X^{HillSlope} + EC50^{HillSlope})
\]

3.6.5 Receptor binding - Key concepts

3.6.5.1 Law of mass action

What is the law of mass action?

Analysis of radioligand binding experiments is based on a simple model, called the law of mass action. This model assumes that binding is reversible.

\[\text{Receptor} + \text{Ligand} \xrightleftharpoons{K_{on}} K_{off} \text{Receptor} \cdot \text{Ligand}\]
Binding occurs when ligand and receptor collide due to diffusion, and when the collision has the correct orientation and enough energy. The rate of association is:

\[
\text{Number of binding events per unit of time} = [\text{Ligand}] \times [\text{Receptor}] \times k_{\text{on}}.
\]

Once binding has occurred, the ligand and receptor remain bound together for a random amount of time. The probability of dissociation is the same at every instant of time. The receptor doesn't "know" how long it has been bound to the ligand. The rate of dissociation is:

\[
\text{Number of dissociation events per unit time} = [\text{ligand} \times \text{receptor}] \times k_{\text{off}}.
\]

After dissociation, the ligand and receptor are the same as at they were before binding. If either the ligand or receptor is chemically modified, then the binding does not follow the law of mass action.

Equilibrium is reached when the rate at which new ligand×receptor complexes are formed equals the rate at which the ligand×receptor complexes dissociate. At equilibrium:

\[
[\text{Ligand}] \cdot [\text{Receptor}] \cdot k_{\text{on}} = [\text{Ligand} \cdot \text{Receptor}] \cdot k_{\text{off}}.
\]

**Meaning of Kd**

Rearrange that equation to define the equilibrium dissociation constant Kd.

\[
\frac{[\text{Ligand}] \cdot [\text{Receptor}]}{[\text{Ligand} \cdot \text{Receptor}]} = \frac{k_{\text{off}}}{k_{\text{on}}} = K_d.
\]

The Kd has a meaning that is easy to understand. Set [Ligand] equal to Kd in the equation above. The Kd terms cancel out, and you will see that [Receptor]/[Ligand×Receptor]=1, so [Receptor] equals [Ligand×Receptor]. Since all the receptors are either free or bound to ligand, this means that half the receptors are free and half are bound to ligand. In other words, when the concentration of ligand equals the Kd, half the receptors will be occupied at equilibrium. If the receptors have a high affinity for the ligand, the Kd will be low, as it will take a low concentration of ligand to bind half the receptors.
The term "dissociation constant"

Don't mix up Kd, the equilibrium dissociation constant, with koff, the dissociation rate constant. They are not the same, and aren't even expressed in the same units.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Name</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>kon</td>
<td>Association rate constant or on-rate constant</td>
<td>M⁻¹ min⁻¹</td>
</tr>
<tr>
<td>koff</td>
<td>Dissociation rate constant or off-rate constant</td>
<td>min⁻¹</td>
</tr>
<tr>
<td>Kd</td>
<td>Equilibrium dissociation constant</td>
<td>M</td>
</tr>
</tbody>
</table>

Fractional occupancy

The law of mass action predicts the fractional receptor occupancy at equilibrium as a function of ligand concentration. Fractional occupancy is the fraction of all receptors that are bound to ligand.

\[
\text{Fractional occupancy} = \frac{[\text{Ligand} \cdot \text{Receptor}]}{[\text{Total Receptor}]} = \frac{[\text{Ligand} \cdot \text{Receptor}]}{[\text{Receptor}] + [\text{Ligand} \cdot \text{Receptor}]} 
\]

This equation is not useful, because you don't know the concentration of unoccupied receptor, [Receptor]. A bit of algebra creates a useful equation.

\[
\text{Fractional occupancy} = \frac{[\text{Ligand}]}{[\text{Ligand}] + K_d} 
\]

This equation assumes equilibrium. To make sense of it, think about a few different values for [Ligand].

<table>
<thead>
<tr>
<th>[Ligand]</th>
<th>Fractional Occupancy</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0%</td>
</tr>
<tr>
<td>1.Kd</td>
<td>50%</td>
</tr>
<tr>
<td>4.Kd</td>
<td>80%</td>
</tr>
<tr>
<td>9.Kd</td>
<td>90%</td>
</tr>
<tr>
<td>99.Kd</td>
<td>99%</td>
</tr>
</tbody>
</table>
Note that when [Ligand]=Kd, fractional occupancy is 50%.

**Assumptions**

Although termed a "law", the law of mass action is simply a model that can be used to explain some experimental data. Because it is so simple, the model is not useful in all situations. The model assumes:

- All receptors are equally accessible to ligands.
- Receptors are either free or bound to ligand. It doesn't allow for more than one affinity state, or states of partial binding.
- Binding does not alter the ligand or receptor.
- Binding is reversible.

Despite its simplicity, the law of mass action has proven to be very useful in describing many aspects of receptor pharmacology and physiology.

### 3.6.5.2 Nonspecific binding

In addition to binding to receptors of interest, radioligands also bind to other sites. Binding to the receptor of interest is called specific binding, while binding to the other sites is called nonspecific binding. This means that nonspecific binding can represent several phenomena:

- In most cases, the bulk of nonspecific binding represents some sort of interaction of the ligand with membranes. The molecular details are unclear, but nonspecific binding depends on the charge and hydrophobicity of a ligand – but not its exact structure.
• Nonspecific binding can also be binding to receptors, transporters, or other proteins not of interest to the investigator. For example binding of the adrenoceptor agonist, epinephrine, to serotonin receptors or metabolic enzymes can be considered “nonspecific”.

• Nonspecific binding can also be binding to the filters used to separate bound from free ligand.

Nonspecific binding is usually (but not necessarily) proportional to the concentration of radioligand (within the range it is used). Add twice as much radioligand, and you will see twice as much nonspecific binding.

Nonspecific binding is detected by measuring radioligand binding in the presence of a saturating concentration of an unlabeled drug that binds to the receptors. Under those conditions, virtually all the receptors are occupied by the unlabeled drug so the radioligand can only bind to nonspecific sites. Subtract the nonspecific binding at a particular concentration of radioligand from the total binding at that concentration to calculate the specific radioligand binding to receptors.

Which unlabeled drug should you use for determining nonspecific binding? The obvious answer is to use the same compound as the radioligand, but in its unlabeled form. In many cases this is necessary, as no other drug is known to bind to the receptors. But most investigators avoid using the same compound as the hot and cold ligand and prefer to define nonspecific binding with a drug that is chemically distinct from the radioligand but which binds to the same receptor.

What concentration of unlabeled drug should you use? You want to use enough to block virtually all the specific radioligand binding, but not so much that you cause more general physical changes to the membrane that might alter binding. If you are studying a well-characterized receptor, a useful rule-of-thumb is to use the unlabeled compound at a concentration equal to 100 times its Kd for the receptors, or 100 times the highest concentration of radioligand, whichever is higher.

Ideally, you should get the same results defining nonspecific binding with a range of concentrations of several drugs, and you should test this when possible. In many assay systems, nonspecific binding is only 10-20% of the total radioligand binding. If the nonspecific binding makes up more than half of the total binding, you will find it hard to get quality data. If your system has a lot of nonspecific binding, try different kinds of filters,
a larger volume of washing buffer, warmer washing buffer, or a different radioligand.

3.6.5.3 Ligand depletion

In many experimental situations, you can assume that a very small fraction of the ligand binds to receptors (or to nonspecific sites). In these situations, you can also assume that the free concentration of ligand is approximately equal to the concentration you added. This assumption vastly simplifies the analysis of binding experiments, and the standard analysis methods depend on this assumption.

In other situations, a large fraction of the ligand binds to the receptors (or binds nonspecifically). This means that the concentration of ligand free in the solution does not equal the concentration you added. The discrepancy is not the same in all tubes or at all times. The free ligand concentration is depleted by binding.

Many investigators use this rule of thumb: If less than 10% of the ligand binds, don't worry about ligand depletion; if more than 10% of the ligand binds, you have three choices:

- Change the experimental conditions. Increase the reaction volume without changing the amount of tissue. The problem with this approach is that it requires more radioligand, which is usually very expensive.

- Measure the free concentration of ligand in every tube. This is possible if you use centrifugation or equilibrium dialysis, but is quite difficult if you use vacuum filtration to remove free radioligand.

- Use analysis techniques that adjust for the difference between the concentration of added ligand and the concentration of free ligand. Prism includes such models for analyzing saturation and competition data. These special analyses only work with radioactive ligands, so the assessment of added ligand and bound ligand are in the same counts-per-minute units. These methods don't work with fluorescent ligands.

3.6.5.4 The radioactivity web calculator

GraphPad Software provides a free radioactivity calculator on
graphpad.com. Use it to perform seven common calculations.

<table>
<thead>
<tr>
<th>Calculation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isotope decay</td>
<td>Calculates radioactive decay during a specified number of days. Select one of the common isotopes, or enter the half-life of another isotope.</td>
</tr>
<tr>
<td>Conc. of stock</td>
<td>Enter mCi/ml and Ci/mmole, which should be on the label. If you are using a molecule labeled with 125I, the specific activity equals 2200 Ci/mmole if each molecule is labeled with one iodine. Also enter the percent of the original isotope remaining (calculated above). The calculations assume that the decay product is not biologically active, so the concentration of stock that is biologically active decreases over time.</td>
</tr>
<tr>
<td>Dilution of stock</td>
<td>Enter the concentration in your stock solution, after accounting for decay. Also enter the concentration and volume you want. The result is the volume of stock you need to use.</td>
</tr>
<tr>
<td>Specific activity (cpm/fmol)</td>
<td>Enter the specific radioactivity as Ci/mmol which should be on the label. If you are using a molecule labeled with 125I, the specific activity equals 2200 Ci/mmol if each molecule is labeled with one iodine. Also enter the counter efficiency - the fraction of radioactive disintegrations that are detected. The efficiency depends on the isotope and instrumentation. With low energy isotopes such as tritium, the efficiency also depends on the experimental details such as the choice of scintillation fluid, the amount of water in the sample, and the presence of any colored substances in the sample.</td>
</tr>
<tr>
<td>Cpm to fmol/mg</td>
<td>Enter the specific radioactivity as cpm/fmol, the number of cpm counted, and the protein content</td>
</tr>
</tbody>
</table>
### Calculation Description

- **of the sample in mg. The result is the number of binding sites in fmol/mg protein.**

- **Cpm to sites/cell** Enter the specific radioactivity as cpm/fmol, the number of cpm counted, and the cell count. The result is the number of binding sites per cell.

- **Cpm to nM** Enter the specific radioactivity as cpm/fmol, the number of cpm counted, and the volume counted. The result is the concentration of radioligand in nM.

---

#### 3.6.6 Receptor binding - Saturation binding

#### 3.6.6.1 Key concepts: Saturation binding

**What is saturation binding?**

In a saturation binding experiment, you vary the concentration of radioligand and measure binding at equilibrium. The goal is to determine the $K_d$ (ligand concentration that binds to half the receptor sites at equilibrium) and $B_{\text{max}}$ (maximum number of binding sites).

**Total, nonspecific and specific binding**

The ligand binds not only to receptors sites, but also to nonspecific sites. There are three approaches to dealing with nonspecific binding.

- Subtract off the nonspecific, and analyze only the specific binding.

- **Analyze the total binding only**, inferring the amount of nonspecific binding from the shape of the total binding curve.

- Globally fit total and nonspecific binding together.

We recommend the third approach (global fitting of total and nonspecific). The problem with fitting specific binding is that you have to make some assumptions in order to subtract nonspecific from total, and the resulting values that you fit aren't really data. When possible, we suggest that you fit the data you actually collect, and avoid creating derived data sets (specific binding, in this case).
Fitting total binding only requires less data, so saves experimental time and money. But most people feel unconformable defining nonspecific binding purely from the shape of a binding curve, without experimentally measuring nonspecific binding. One advantage of fitting total binding only is that equations have been derived for fitting such data, even when a substantial fraction of the ligand binds, resulting in ligand depletion (free concentration substantially less than the added concentration).

**One-vs. two sites**

Prism offers models for fitting one or two sites. You can use choices in the Compare tab to compare the two fits. When comparing the fits to the one- and two-site models, use common sense as well as statistics. Don't accept a two site model, if one of the sites is only a tiny fraction of the total, or if its Kd is outside the range of radioligand concentrations you used in the experiment.

### 3.6.6.2 Equation: One site -- Total binding

**Introduction**

You don't have to measure nonspecific binding directly. Instead, you can determine Bmax and Kd by fitting only total binding by assuming that the amount of nonspecific binding is proportional to the concentration of radioligand.

**Step by step**

Create an XY data table. Enter radioligand concentration into X, and total binding into Y. If you have several experimental conditions, place the first into column A, the second into column B, etc.

Use any convenient units for X and Y. The Kd will be reported in the same units as X, and the Bmax will be reported in the same units as Y.

From the table of total binding, click Analyze, choose nonlinear regression, choose the panel of Saturation Binding equations, and choose *One site -- Total.*
Consider constraining the parameter Background to a constant value of zero. This is the measured 'binding' when there is no radioligand binding added, so represents the counter background, if there is any.

**Model**

\[ Y = \frac{B_{\text{max}} \cdot X}{(K_d + X)} + NS \cdot X + \text{Background} \]

**Interpret the parameters**

- **Bmax** is the maximum specific binding in the same units as Y.
- **Kd** is the equilibrium binding constant, in the same units as X. It is the radioligand concentration needed to achieve a half-maximum binding at equilibrium.
- **NS** is the slope of nonspecific binding in Y units divided by X units.
- **Background** is the amount of nonspecific binding with no added radioligand. This represents counter background. If your counter automatically subtracts off the background signal, you can constrain Background to a constant value of zero.

**Notes**

This analysis assumes that only a small fraction of radioligand binds, which means that the concentration you added is virtually identical to the free concentration. If you can't make this assumption, use an alternative analysis.
3.6.6.3 Equation: One site -- Fit total and nonspecific binding

Introduction

In a saturation binding experiment, you vary the concentration of radioligand and measure binding. The goal is to determine the Kd (ligand concentration that binds to half the receptor sites at equilibrium) and Bmax (maximum number of binding sites).

The ligand binds not only to receptors sites, but also to nonspecific sites. There are three approaches to dealing with nonspecific binding.

- Subtract off the nonspecific, and analyze only the specific binding.
- Analyze the total binding only, inferring the amount of nonspecific binding from the shape of the total binding curve.
- Globally analyze the total and nonspecific binding at one time. This is the best approach, and the details are explained below.

Step by step

Create an XY data table. Enter radioligand concentration into X, total binding into Y, and nonspecific binding into column B.

Use any convenient units for X. The Kd will be reported in those same concentration units. Use the same units for total and nonspecific binding. The Bmax will be reported in those same units.

Alternatively choose the sample data set: Binding - Saturation binding to total and nonspecific.

From the data table, click Analyze, choose nonlinear regression, choose the panel of Saturation Binding equations, and choose One site -- Total and nonspecific binding.

Consider constraining the parameter Background to a constant value of zero. This is the measured 'binding' when there is no radioligand binding added, so represents the counter background, if there is any.

Model

\[ \text{specific} = \frac{\text{Bmax} \times X}{X + \text{Kd}} \]
\[ \text{nonspecific} = \text{NS} \times X + \text{Background} \]
\[ Y = \text{specific} + \text{nonspecific} \]
The \(<A>Y=nonspecific</A>\) syntax means that the third line is only used for data set A (total binding) while the fourth line is used only for data set B (nonspecific).

The parameters NS and Background are shared between the two data sets.

**Interpret the parameters**

- **Bmax** is the maximum specific binding in the same units as Y. It is the specific binding extrapolated to very high concentrations of radioligand, so its value is almost always higher than any specific binding measured in your experiment.

- **Kd** is the equilibrium binding constant, in the same units as X. It is the radioligand concentration needed to achieve a half-maximum binding at equilibrium.

- **NS** is the slope of nonspecific binding in Y units divided by X units.

- **Background** is the amount of nonspecific binding with no added radioligand. This represents counter background. If your counter automatically subtracts off the background signal, you can constrain Background to a constant value of zero.
3.6.6.4 Equation: One site -- Total, accounting for ligand depletion

**Introduction**

You don't have to measure nonspecific binding directly. Instead, you can determine Bmax and Kd by fitting only total binding by assuming that the amount of nonspecific binding is proportional to the concentration of radioligand.

If only a small fraction of radioligand binds, you can use a simpler model. This equation allows for a substantial fraction of the added ligand to bind. This only works with radioactive ligands, so the assessment of added ligand and bound ligand are in the same counts-per-minute units. This method doesn't work with fluorescent ligands.

**Step by step**

Create an XY data table. Enter radioligand concentration into X, and total binding into Y. If you have several experimental conditions, place the first into column A, the second into column B, etc.

Enter both X and Y in CPM units. This is essential for the analysis to work.

From the table of total binding, click Analyze, choose nonlinear regression, choose the panel of Saturation Binding equations, and choose One site -- Total, accounting for ligand depletion.

You must constrain two parameters to constant values based on your experimental design:

- SpAct is the specific radioactivity in cpm/fmol
- Vol is the reaction volume in ml

**Model**

\[
K_{dCPM} = K_d M \cdot V o l \cdot 1000 \cdot S p e c A c t \]

\[
a = -1 - N S
\]

\[
b = K_{dCPM} + N S \cdot K_{dCPM} + X + 2 \cdot X \cdot N S + B_{max}
\]

\[
c = -1 \cdot X \cdot (N S \cdot K_{dCPM} + X \cdot N S + B_{max})
\]

\[
Y = \frac{-b + \sqrt{b^2 - 4 \cdot a \cdot c}}{2 \cdot a} \quad ; Y \ is \ in \ cpm
\]
Interpret the parameters

**Bmax** is the maximum specific binding in cpm.

**KdnM** is the equilibrium binding constant in nM.

**NS** is the slope of nonspecific binding in Y units divided by X units.

Notes

This analysis accounts for the fact that a large fraction of the added radioligand binds to the receptors. If you are able to assume that only a small fraction of radioligand binds, which means that the concentration you added is virtually identical to the free concentration, use an alternative analysis.

Reference

This equation came from S. Swillens (Molecular Pharmacology, 47: 1197-1203, 1995)

3.6.6.5  Equation: One site -- Specific binding

**Introduction**

In a saturation binding experiment, you vary the concentration of radioligand and measure binding. The goal is to determine the Kd (ligand concentration that binds to half the receptor sites at equilibrium) and Bmax (maximum number of binding sites).
The ligand binds not only to receptors sites, but also to nonspecific sites. There are three approaches to dealing with nonspecific binding.

- Subtract off the nonspecific, and analyze only the specific binding. Read on for this approach.
- Analyze the total binding only, inferring the amount of nonspecific binding from the shape of the total binding curve. Learn more.
- Globally analyze the total and nonspecific binding at one time. Learn more.

**Step by step**

Create an XY data table. Enter radioligand concentration into X, and specific binding into Y. If you have several experimental conditions, place the first into column A, the second into column B, etc.

An alternative approach would be to enter total binding into column A, and nonspecific into column B. Then use the Remove Baseline analysis to subtract column B from column A, creating a new results table with the specific binding.

From the table of specific binding, click Analyze, choose nonlinear regression, choose the panel of Saturation Binding equations, and choose *One site specific binding*.

**Model**

\[ Y = \frac{B_{\text{max}} \times X}{K_d + X} \]
Interpret the parameters

**B\text{max}** is the maximum specific binding in the same units as Y. It is the specific binding extrapolated to very high concentrations of radioligand, and so its value is almost always higher than any specific binding measured in your experiment.

**K\text{d}** is the equilibrium binding constant, in the same units as X. It is the radioligand concentration needed to achieve a half-maximum binding at equilibrium.

Create a Scatchard plot

Before nonlinear regression was available, investigators had to transform curved data into straight lines, so they could analyze with linear regression. One way to do this is with a Scatchard plot, which plots specific binding vs. the ratio of specific binding to the concentration of free radioligand.

If you create a Scatchard plot, use it only to display your data. The human retina and visual cortex evolved to detect edges (straight lines),
not rectangular hyperbolas, and so it can help to display data this way. Scatchard plots are often shown as insets to the saturation binding curves. They are especially useful when you want to show a change in $B_{max}$ or $K_d$.

Don't use the slope and intercept of a linear regression line to determine values for $B_{max}$ and $K_d$. If you do this, you won't get the most accurate values for $B_{max}$ and $K_d$. The problem is that the transformation distorts the experimental error, so the data on the Scatchard plot do not obey the assumptions of linear regression. Use nonlinear regression to obtain the most accurate values of $K_d$ and $B_{max}$.

To create a Scatchard plot from your specific binding data, use Prism's Transform analysis, and choose the Scatchard transform from the panel of biochemistry and pharmacology transforms.

To create a Scatchard line corresponding to the nonlinear regression fit, follow these steps:

1. Create a new XY data table, with no subcolumns.

2. Into row 1 enter $X=0$, $Y=B_{max}/K_d$ (previously determined by nonlinear regression). You need to do the calculation manually, and enter a number.

3. Into row 2 enter $X=B_{max}$ and $Y=0$. Again enter the number into the $X$ column, not the text 'Bmax'.

4. Note the name of this data table. Perhaps rename it to something appropriate.

5. Go to the Scatchard graph.

6. Drag the new table from the navigator and drop onto the graph.

7. Double-click on one of the new symbols for that data set to bring up the Format Graph dialog.

8. Choose to plot no symbols, but to connect with a line.
Notes

- This is not the best way to determine $B_{\text{max}}$ and $K_d$. It is better to **globally fit total and nonspecific binding**[^1], without subtracting to compute specific binding.

- When making a Scatchard plot, you have to choose what units you want to use for the Y-axis. Some investigators express both free ligand and specific binding in cpm so the ratio bound/free is a unitless fraction. While this is easy to interpret (it is the fraction of radioligand bound to receptors), an alternative is to express specific binding in sites/cell or fmol/mg protein, and to express the free radioligand concentration in nM. While this makes the Y-axis hard to interpret visually, it provides correct units for the slope (which equals $-1/K_d$).

3.6.6.6 Binding potential

**The problem: Binding isotherms without going to high enough concentrations**

![Binding isotherms graph](image)

The figure above shows a saturation binding equation fit to specific binding data. But all the concentrations are relatively low (compared to the $K_d$ of binding) so the data almost form a straight line. The saturation binding model fits the data fine (solid curve), with a narrow 95% confidence band around the curve (dashed lines). But look at the confidence interval for $B_{\text{max}}$ and $K_d$ (shown in the box)! They are super wide, even descending into negative (impossible) values. With these

[^1]: Source: [GraphPad Software](https://www.graphpad.com)
data, the curve fit the data fine, but neither parameter in the model (Bmax and Kd) were fit with reasonable confidence intervals. This is no surprise. The data don’t show even a hint of plateauing, so the data simply don’t define the Bmax and Kd at all.

The problem is that the data are consistent with a system with a huge number of low affinity receptors (high Bmax; high Kd) or a smaller number of high affinity receptors (low Bmax; low Kd). The figure below shows two ways the visualize this. The graph on the left shows the same fit as the one above, but with the best-fit curve and its confidence bands extended out to larger concentrations. While the confidence band is tight near the points, is it super wide as it goes beyond concentrations with data. The graph on the right show two different fits, with the Kd constrained to equal 200 nM or 500 nM. The curves are very different, yet both go near the data.

Another way to look at the problem with these data is that the Bmax and Kd are correlated. Prism can report the covariance matrix as part of its nonlinear regression results. With only two parameters, there is only one value in that "matrix". The covariance between Kd and Bmax (which can range from 0.0 to 1.0) is 0.9993. Prism can also report the dependency of each parameter (which also can range from 0.0 to 1.0). With only two parameters, both have the same dependency, which is 0.9986 for this example. This is not quite high enough for Prism to declare the results ambiguous, but the threshold for that designation (dependency > 0.9999) is arbitrary.

It is tempting to give up at this point and say that nothing can be determined without more data at higher concentrations. But in some systems, especially those using PET scanning to detect receptors, data
like this are typical. It is impossible to use higher concentrations of ligand.

**Fitting the binding potential**

What can be learned from these data? The data are nearly linear. What does the slope of that line tell us? The slope, or rather the initial slope, is termed the *binding potential* (1). Don’t be misled by the term potential. It has nothing to do with potential energy and thermodynamics. Rather it is a measure of how much binding will occur with a small amount of ligand.

If you start with the specific binding equation, take its derivative with respect to $X$ (which is concentration), and set $X$ equal to zero, you’ll derive the initial slope of a specific binding curve. It turns out that this slope, the binding potential, equals $B_{max}/K_d$. $B_{max}$ is measured in the same units as the $Y$ values in the data. $K_d$ is measured in the same units as the $X$ values. So the binding potential has units equal to the $Y$ units divided by the $X$ units.

Prism can fit a specific binding curve, and also report the ratio of $B_{max}/K_d$ with its confidence interval. To do this either create a new user-defined equation or (easier) clone the built in one. Then go to the last tab of the dialog that defines the equation. “Transforms to Report”. Choose to report the ratio of two parameters, and define the numerator and denominator.
In this example, Prism finds that the Binding Potential is 242.1, with a confidence interval ranging from 183.4 to 300.7. That confidence interval is reasonably narrow, so the result is quite useful.

**Reference**


[Download the Prism file for this example.](#)

### 3.6.6.7 Equation: Two sites -- Specific binding only

**Introduction**

In a saturation binding experiment, you vary the concentration of radioligand and measure binding. The goal is to determine the Kd (ligand concentration that binds to half the receptor sites at equilibrium) and Bmax (maximum number of binding sites) of both kinds of receptors.
The ligand binds not only to receptors sites, but also to nonspecific sites. There are three approaches to dealing with nonspecific binding.

- Subtract off the nonspecific, and analyze only the specific binding. Read on for this approach.

- Analyze the total binding only, inferring the amount of nonspecific binding from the shape of the total binding curve. This approach doesn't work well when there are two classes of receptors.

- Globally analyze the total and nonspecific binding at one time. Learn more.

**Step by step**

Create an XY data table. Enter radioligand concentration into X, and specific binding into Y. If you have several experimental conditions, place the first into column A, the second into column B, etc.

An alternative approach would be to enter total binding into column A, and nonspecific into column B. Then use the Remove Baseline analysis to subtract column B from column A, creating a new results table with the specific binding.

From the table of specific binding, click Analyze, choose nonlinear regression, choose the panel of Saturation Binding equations, and choose *Two sites -- Specific binding.*

**Model**

\[
\begin{align*}
\text{Site1} &= \frac{B_{max Hi} \times X}{K_{d Hi} + X} \\
\text{Site2} &= \frac{B_{max Lo} \times X}{K_{d Lo} + X} \\
Y &= \text{Site1} + \text{Site2}
\end{align*}
\]
Interpret the parameters

\[ \text{BmaxHi} \] and \[ \text{BmaxLo} \] are the maximum specific bindings to the two sites in the same units as Y.

\[ \text{KdHi} \] and \[ \text{KdLo} \] are the equilibrium binding constants, in the same units as X. It is the radioligand concentration needed to achieve a half-maximum binding at equilibrium.

Scatchard plots of two site binding

The left panel below shows binding of a radioligand to two independent binding sites present in equal concentrations, but with a tenfold difference in \[ \text{Kd} \]. The two individual curves are shown as dotted and dashed curves. When you do the experiment, you can't observe the individual components, but observe the sum, which is shown as a solid curve. Note that this curve is not obviously biphasic.

The right panel shows the same data plotted on a Scatchard plot. The binding to each receptor is shown as a straight line (dotted, or dashed). The total binding, plotted on a Scatchard plot, is curved. Note that the two lines that represent binding to each type of receptor are NOT the asymptotes of the curve.
To plot the two straight lines that correspond to the nonlinear regression fit, create a new data table that defines the two lines as shown below, using Bmax and Kd values determined by nonlinear regression.

<table>
<thead>
<tr>
<th>X</th>
<th>A</th>
<th>B</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Bmax1/Kd1</td>
<td>0</td>
</tr>
<tr>
<td>Bmax1</td>
<td>0</td>
<td>Bmax2/Kd2</td>
</tr>
<tr>
<td>0</td>
<td>Bmax2/Kd2</td>
<td>0</td>
</tr>
</tbody>
</table>

Go to the graph of the Scatchard transformed data and drag the new table to that graph. Use the Format Graph dialog to plot the two datasets from the table using connecting lines but no symbols.

3.6.6.8 Equation: Two sites -- Fit total and nonspecific binding

Introduction

In a saturation binding experiment, you vary the concentration of radioligand and measure binding. The goal is to determine the Kd (ligand concentration that binds to half the receptor sites at equilibrium) and Bmax (maximum number of binding sites).

The ligand binds not only to receptors sites, but also to nonspecific sites. There are three approaches to dealing with nonspecific binding.

- Subtract off the nonspecific, and analyze only the specific binding.
- Analyze the total binding only, inferring the amount of nonspecific binding from the shape of the total binding curve. This approach doesn't work well when the ligand binds to two sites.
- Globally analyze the total and nonspecific binding at one time. This is the best approach, and the details are explained below.
Step by step

Create an XY data table. Enter radioligand concentration into X, total binding into Y, and nonspecific binding into column B.

Use any convenient units for X. The Kd will be reported in those same concentration units. Use the same units for total and nonspecific binding. The Bmax will be reported in those same units.

From the data table, click Analyze, choose nonlinear regression, choose the panel of Saturation Binding equations, and choose *Two sites -- Total and nonspecific binding*.

The parameter Background is the measured 'binding' when there is no radioligand binding added, so represents the counter background, if there is any. Consider constraining it to a constant value of zero.

Model

\[
\begin{align*}
\text{Specific1} &= \text{BmaxHi} \times X \div (X + \text{KdHi}) \\
\text{Specific2} &= \text{BmaxLo} \times X \div (X + \text{KdLo}) \\
\text{Nonspecific} &= \text{NS} \times X + \text{Background} \\
\text{<A>Y} &= \text{Specific1} + \text{Specific2} + \text{Nonspecific} \\
\text{<B>Y} &= \text{Nonspecific}
\end{align*}
\]

The <A> and <B> syntax means that the fourth line is only used for data set A (total binding) while the fifth line is used only for data set B (nonspecific).

The parameters NS and Background are shared between the two data sets.

Interpret the parameters

**BmaxHi** and **BmaxLo** are the maximum specific bindings to the two sites in the same units as Y.

**KdHi** and **KdLo** are the equilibrium binding constants, in the same units as X. It is the radioligand concentration needed to achieve a half-maximum binding at equilibrium.

**NS** is the slope of nonspecific binding in Y units divided by X units.
Background is the amount of nonspecific binding with no added radioligand. This represents counter background. If your counter automatically subtracts off the background signal, you can constrain Background to a constant value of zero.

3.6.6.9 Equation: One site with allosteric modulator

Introduction

An allosteric modulator can reduce radioligand binding. This model fits experiments, where entire radioligand binding curves are measured in the absence and presence of modulator. The goal is to learn the affinity of the modulator for binding to its site, and also determine the value of alpha, the ternary complex constant that quantifies the degree to which binding of the modulator alters the affinity of the radioligand for the receptor site.

Step by step

Create an XY data table. Enter the concentration of the labeled ligand into X, using any convenient units (maybe nM). Enter specific binding into Y in any convenient units. Enter data with no modulator into column A. Enter data collected with a constant concentration of modulator into column B. Repeat, if you have data, for column C, D, E, ..., each with a different concentration of modulator. Enter the modulator concentration (in nanomolar so 1nM is entered as '1') into the column titles. Don't forget to enter '0' as the column title for data set A.

From the data table, click Analyze, choose nonlinear regression, choose the panel of Saturation Binding equations, and choose Allosteric modulator shift.

You don't need to constrain any parameters to constant values.

Model

Hot=X
Alpha=10^logalpha
KB=10^logKB
KApp=KDHot*((1+Allo/KB)/(1+alpha*Allo/KB))
Y=Bmax*Hot/(Hot+KApp)
Interpret the parameters

**$K_b$** is the equilibrium dissociation constant (Molar) of modulator binding to its allosteric site. It is in the same molar units used to enter the modulator concentration into column titles on the data table.

**$K_{dHOT}$** is the equilibrium dissociation constant of the radioligand. It is expressed in the same units used to enter X values, nM in the example.

**Alpha** is the ternary complex constant. When alpha=1.0, the modulator won't alter binding. If alpha is less than 1.0, then the modulator reduces ligand binding. If alpha is greater than 1.0, then the modulator increases binding. In the example shown about, alpha equals 0.01 so the modulator greatly decreases binding.

Notes

- This model is designed to analyze data when the unlabeled compound works via an allosteric site. Since the labeled and unlabeled ligands are acting via different sites, it is inappropriate (and incorrect) to refer to the modulator as a competitor.

- The model is written to fit the logarithm of alpha, rather than alpha itself. This is because alpha is asymmetrically (all values from 0 to 1 mean that the modulator decreases binding, while all values from 1 to infinity mean that the modulator enhances binding. On a log scale, its values are more symmetrical, so the confidence interval computed on a log scale (as Prism does) are more accurate. Prism reports both alpha and log(alpha).
• This model assumes that the allosteric modulator is present in excess, so the concentration you added is very close to its free concentration. This model won't work when the concentration of allosteric modulator is limiting (as it is when G proteins alter agonist binding to many receptors). No explicit model can handle this situation. You need to define the model with an implicit equation (Y on both sides of the equals sign) and Prism cannot handle such equations.

Reference


3.6.6.10 Equation: Specific binding with Hill slope

Introduction

In a saturation binding experiment, you vary the concentration of radioligand and measure binding. The goal is to determine the Kd (ligand concentration that binds to half the receptor sites at equilibrium) and Bmax (maximum number of binding sites).

This equation assumes you have subtracted off the nonspecific, and are only analyzing specific binding.

This equation fits a Hill slope. If you assume the Hill slope is 1.0 (for mass action binding of a monomer to one site) use a simpler equation. 

Step by step

Create an XY data table. Enter radioligand concentration into X, and specific binding into Y. If you have several experimental conditions, place the first into column A, the second into column B, etc.

An alternative approach would be to enter total binding into column A, and nonspecific into column B. Then use the Remove Baseline analysis to subtract column B from column A, creating a new results table with the specific binding.

From the table of specific binding, click Analyze, choose nonlinear regression, choose the panel of Saturation Binding equations, and choose Specific binding with Hill Slope.
Model

\[ Y = \frac{B_{\text{max}} \cdot X^h}{(K_d^h + X^h)} \]

Interpret the parameters

**B_{\text{max}}** is the maximum specific binding in the same units as Y. It is the specific binding extrapolated to very high concentrations of radioligand, and so its value is almost always higher than any specific binding measured in your experiment.

**K_d** is the radioligand concentration needed to achieve a half-maximum binding at equilibrium, expressed in the same units as X. If \( h = 1.0 \), this is the equilibrium binding constant. If \( h \) is not equal to 1.0, then the molecular interpretation of the K_d depends on why \( h \) is not 1.0.

**h** is the Hill slope. It equals 1.0 when a monomer binds with no cooperativity to one site. When it is greater than 1.0, you see a sigmoidal look to the graph as shown above. This happens when the receptor or ligand has multiple binding sites with positive cooperativity. The Hill slope is less than zero when there are multiple binding sites with different affinities for ligand or when there is negative cooperativity.

Note

Before version 7, this equation was called "One site -- specific binding with Hill slope". We removed "one site" from the name because if the Hill slope is not 1.0, this equation doesn't really describe binding to one site.
3.6.7 Receptor binding - Competitive binding

3.6.7.1 Key concepts: Competitive binding

**What is competitive binding?**

In a competitive binding experiment, you use a single concentration of labeled (hot) ligand and vary the concentration of unlabeled (cold) drugs, and measure binding at equilibrium.

**Comparing one- and two-site models**

Prism offers models for fitting one or two sites. You can use choices in the Compare tab to compare the two fits. When comparing the fits to the one- and two-site models, use common sense as well as statistics. Don't accept a two site model, if one of the sites is only a tiny fraction of the total, or if its Ki is outside the range of competitor concentrations you used in the experiment.

**Ligand depletion**

If a large fraction of the added radioligand binds to the receptors, the ligand is depleted so the concentration you added is greater than the free concentration. You need to fit these data to a model that accounts for ligand depletion.

**Homologous binding**

An homologous binding experiment is one where the labeled and unlabeled ligands have identical affinities for the receptors. Generally this is because the two are chemically identical. Receptor number and affinity are determined by analyzing the competition of varying concentrations of unlabeled ligand for one (or better, two) concentrations of labeled ligand. Prism offers a special model for fitting homologous competition experiments.

**Allosteric modulators**

Allosteric modulators can alter radioligand binding, even though they bind to different sites. Since the hot and cold ligands bind to different sites, the term 'competition' is not apt, but we include this model here.
because the experimental design is the same as used for competitive binding. Prism can fit binding inhibition (or augmentation) by an allosteric modulator based on the ternary complex model. Note that this model assumes the allosteric modulator is present in excess, so is not depleted by binding to the receptors.

### 3.6.7.2 Equation: One site - Fit Ki

**Introduction**

You can determine the equilibrium dissociation constant of an unlabelled ligand by measuring its competition for radioligand binding.

**Step by step**

Create an XY data table. Enter the logarithm of the concentration of the unlabeled compound into X, and binding into Y. If you have several experimental conditions, place the first into column A, the second into column B, etc. Use subcolumns to enter replicates.

From the data table, click Analyze, choose nonlinear regression, choose the panel of Competition Binding equations, and choose *One site - Fit Ki*.

You must constrain two parameters to constant values based on your experimental design:

- RadioligandNM is the concentration of labeled ligand in nM. A single concentration of radioligand is used for the entire experiment.

- HotKdNM is the equilibrium dissociation constant of the labeled ligand in nM.

**Model**

$$\log EC50 = \log(10^{\log Ki} \cdot (1 + \text{RadioligandNM}/\text{HotKdNM}))$$

$$Y = \text{Bottom} + \frac{(\text{Top} - \text{Bottom})}{1 + 10^{X - \log EC50}}$$
Interpret the parameters

Top and Bottom are plateaus in the units of Y axis.

$logKI$ is the log of the molar equilibrium dissociation constant of unlabeled ligand.

$Ki$ is the equilibrium dissociation constant in Molar.

Notes

This model fits the Ki of the unlabelled ligand directly. It does not report the EC50, so you do not need to apply the Cheng and Prusoff correction(1). Instead you enter the concentration of radioligand and its Kd as constants, and Prism directly fits the Ki of your cold compound. If you want to know the IC50, fit a log(dose)-response curve.

If you want to know the IC50 (which is not very informative in this situation) fit the data using an alternative equation\cite{372}.

The analysis assumes that you have one site, and that the binding is reversible and at equilibrium.

3.6.7.3 Equation: One site - Fit logIC50

**Introduction**

You can determine the equilibrium dissociation constant of an unlabelled ligand by measuring its competition for radioligand binding.

**Step by step**

Create an XY data table. Enter the logarithm of the concentration of the unlabeled compound into X, and binding into Y. If you have several experimental conditions, place the first into column A, the second into column B, etc. Use subcolumns to enter replicates.

From the data table, click Analyze, choose nonlinear regression, choose the panel of Competition Binding equations, and choose One site - Fit logEC50.

**Model**

\[
Y = \text{Bottom} + \frac{(\text{Top} - \text{Bottom})}{(1 + 10^{(X - \text{LogIC50})})}
\]

**Interpret the parameters**

**Top** and **Bottom** are plateaus in the units of Y axis.

**logIC50** is the log of the concentration of competitor that results in binding half-way between Bottom and Top.
Notes

This model is the same as an inhibitory dose-response curve. It fits the logIC50, which is not the same as the Ki of the unlabelled ligand for binding. The Ki depends on the IC50, the concentration of radioligand, and its Kd for binding. You can fit the Ki directly using a different equation.[74]

The analysis assumes that you have one site, and that the binding is reversible and at equilibrium.

3.6.7.4 Equation: Two sites - Fit Ki

Introduction

You can determine the equilibrium dissociation constant of an unlabelled ligand by measuring its competition for radioligand binding.

Step by step

Create an XY data table. Enter the logarithm of the Molar concentration of the unlabeled compound into X, and binding into Y. If you have several experimental conditions, place the first into column A, the second into column B, etc. Use subcolumns to enter replicates.

From the data table, click Analyze, choose nonlinear regression, choose the panel of Competition Binding equations, and choose Two sites - Fit Ki.

You must constrain three parameters to constant values based on your experimental design:

- **HotNM** is the concentration of labeled ligand in nM. A single concentration of radioligand is used for the entire experiment.

- **HotKdNMHi** is the equilibrium dissociation constant of the labeled ligand for the high-affinity site in nM.

- **HotKdNMLo** is the equilibrium dissociation constant of the labeled ligand for the low-affinity site in nM.

Model
\[
\text{logEC50Lo} = \text{log}(10^{\text{logKiLo}} \times (1 + \text{HotNM/HotKdNMLo})) \\
\text{logEC50Hi} = \text{log}(10^{\text{logKiHi}} \times (1 + \text{HotNM/HotKdNMHi})) \\
\text{Span} = \text{Top} - \text{Bottom} \\
\text{Part1} = \text{FractionHi} \times \text{Span} / (1 + 10^{(X - \text{LogEC50Hi})}) \\
\text{Part2} = (1 - \text{FractionHi}) \times \text{Span} / (1 + 10^{(X - \text{LogEC50Lo})}) \\
\text{Y} = \text{Bottom} + \text{Part1} + \text{Part2}
\]

**Interpret the parameters**

**Top** and **Bottom** are plateaus in the units of Y axis.

**FractionHi** is the fraction of all the sites that have high affinity for the competitor.

**logKi_Hi** and **logKi_Lo** are the logarithms of the two molar Ki values.

**Notes**

This model fits the two log(Ki) values of the unlabelled ligand directly. It does not report the IC50s, so you do not need to apply the Cheng and Prusoff correction. Instead you enter the concentration of radioligand and its Kd as constants, and Prism directly fits the Ki of your cold compound. If you want to fit the two IC50 values instead of the Ki values, use a different equation.

The analysis assumes that you know the affinity of both sites for the labeled ligand. In many cases, the radioligand has the same affinity for
both sites. In that case, simply enter that value twice. If the two sites have different affinities for the labeled ligand, enter both values (determined from other experiments). Watch out for the labels. The constant \( K_{dHi} \) is the \( K_d \) of the hot ligand for the receptors with the high affinity for the unlabeled ligand, and \( K_{dLo} \) is the \( K_d \) of the hot ligand for the receptors with lower affinity for the unlabeled ligand. So \( K_{dHi} \) may be larger or smaller than \( K_{dLo} \).

This analysis assumes that the binding is reversible and at equilibrium. It also assumes that the labeled and unlabeled ligands compete for the same binding sites.


### 3.6.7.5 Equation: Two sites - Fit logIC50

**Introduction**

You can determine the equilibrium dissociation constant of an unlabelled ligand by measuring its competition for radioligand binding. This model assumes that there are two classes of sites with different affinities for the radioligand and competitor.

**Step by step**

Create an XY data table. Enter the logarithm of the molar concentration of the unlabeled compound into \( X \), and binding into \( Y \). If you have several experimental conditions, place the first into column A, the second into column B, etc. Use subcolumns to enter replicates.

From the data table, click Analyze, choose nonlinear regression, choose the panel of Competition Binding equations, and choose *Two sites - Fit logIC50*.

**Model**

\[
\log EC_{50 Lo} = \log (10^{\log K_{i Lo}} * (1 + HotNM / HotKdNMLo))
\]

\[
\log EC_{50 Hi} = \log (10^{\log K_{i Hi}} * (1 + HotNM / HotKdNMHi))
\]

\[
\text{Span} = \text{Top} - \text{Bottom}
\]

\[
\text{Part1} = \text{FractionHi} * \text{Span} / (1 + 10^{(X - \log EC_{50 Hi})})
\]

\[
\text{Part2} = (1 - \text{FractionHi}) * \text{Span} / (1 + 10^{(X - \log EC_{50 Lo})})
\]

\[
Y = \text{Bottom} + \text{Part1} + \text{Part2}
\]
Interpret the parameters

**Top** and **Bottom** are plateaus in the units of Y axis.

**FractionHi** is the fraction of all the sites that have high affinity for the competitor.

**logIC50Hi** and **logIC50Lo** are the logarithms of the two IC50 values.

Notes

This model fits the two IC50 values of the unlabelled ligand. It does not report the two Ki values. The Ki values depend on the IC50s, the concentration of radioligand, and its Kd for binding. You can fit the Ki values directly using a different equation.

The analysis assumes that you know the affinity of both sites for the labeled ligand. In many cases, the radioligand has the same affinity for both sites. In that case, simply enter that value twice. If the two sites have different affinities for the labeled ligand, enter both values (determined from other experiments). Watch out for the labels. The constant HotKdnmHi is the Kd of the hot ligand for the receptors with the high affinity for the unlabeled ligand, and HotKdnmLo is the Kd of the hot ligand for the receptors with lower affinity for the unlabeled ligand. So HotKdnmHi may be larger or smaller than HotKdnmLo.
This analysis assumes that the binding is reversible and at equilibrium. It also assumes that the labeled and unlabeled ligands compete for the same binding sites.

### 3.6.7.6 Equation: One site - Ligand depletion

#### Introduction

This model for competitive binding is useful when a large fraction of the added radioligand binds to the receptors, so the concentration you added is greater than the free concentration.

These equation allows for a substantial fraction of the added ligand to bind. This only works with radioactive ligands, so the assessment of added ligand and bound ligand are in the same counts-per-minute units. This method doesn't work with fluorescent ligands.

#### Step by step

Create an XY data table. Enter the logarithm of the molar concentration of the unlabeled compound into X, and binding into Y. The Y values must be in cpm.

If you have several experimental conditions, place the first into column A, the second into column B, etc. Use subcolumns to enter replicates.

From the data table, click Analyze, choose nonlinear regression, choose the panel of Competition Binding equations, and choose *One site -- Heterologous with depletion*.

You must constrain four parameters to constant values based on your experimental design:

- **Hot** is the amount of labeled ligand in cpm. A single concentration of radioligand is used for the entire experiment.
- **KdNM** is the equilibrium dissociation constant of the labeled ligand in nM.
- **SpAct** is the specific radioactivity in cpm/fmol.
- **Vol** is the reaction volume in ml.
Model

KdCPM=KdnM*SpAct*vol*1000
; nmol/L *(cpm/fmol * ml * .001L/ml * 1000000fmol/nmol) = cpm
R=NS+1
S=[1+10^(X-LogKi)]*KdCPM+Hot
a=-1*R
b=R*NS+Hot + Bmax
c= -1*Hot*(S*NS + Bmax)
Y= (-1*b + sqrt(b*b-4*a*c))/(2*a)

Interpret the parameters

logKI is the log of the molar equilibrium dissociation constant of unlabeled ligand.

Ki is the equilibrium dissociation constant in Molar.

Bmax is the maximum binding of ligand to receptors in cpm. This represents binding to all the receptors so is higher than the top plateau of the curve.

NS is the fraction of the radioligand that binds to nonspecific sites.

Notes

This analysis accounts for the fact that a large fraction of the added radioligand binds to the receptors. If you are able to assume that only a small fraction of radioligand binds, which means that the concentration
you added is virtually identical to the free concentration, use an alternative analysis.

Reference

This equation came from S. Swillens (Molecular Pharmacology, 47: 1197-1203, 1995)

3.6.7.7 Equation: One site - Homologous

Introduction

An homologous binding experiment is one where the labeled and unlabeled ligands have identical affinities for the receptors. Generally this is because the two are chemically identical. Receptor number and affinity are determined by analyzing the competition of varying concentrations of unlabeled ligand for one (or better, two) concentrations of labeled ligand.

Step by step

Create an XY data table. Enter the logarithm of the concentration of the unlabeled compound into X, and binding into Y. Enter the concentration of labeled ligand (in nM) as the column title. You will get better results if you use two different concentrations of labeled ligand.

If you have several experimental conditions, place the first into column A, the second into column B, etc. Use subcolumns to enter replicates.

From the data table, click Analyze, choose nonlinear regression, choose the panel of Competition Binding equations, and choose One site - Homologous.

Model

ColdNM=10^(x+9) ;Cold concentration in nM
KdNM=10^(logKD+9) ;Kd in nM
Y=(Bmax*HotnM)/(HotnM + ColdNM + KdNM) + Bottom
Interpret the parameters

**logKd** Log of the equilibrium binding constant. Since the analysis is based on the assumption that labeled and unlabeled ligand bind with identical affinities, the logKd applies to both forms of the ligand.

**Bmax** is the maximum binding of ligand to receptors in cpm. This represents binding to all the receptors so is higher than the top plateau of the curve.

**NS** is a measure of nonspecific binding. It is the bottom plateau in units of the Y axis divided by the concentration of the hot ligand in nM. In other words, it is the fraction of hot ligand that binds nonspecifically.

Prism fits the curves globally to find one shared value for logKd, Bmax and NS from all sets of data

**Notes**

This model assumes that the hot and cold ligand binds identically to the receptor, and that you use two concentrations of hot ligand (in column A, and B ..) and vary cold. It assumes that a small fraction of added ligand binds, so the free concentration is close to what you added.
3.6.7.8 Equation: Allosteric modulator

**Introduction**

Fits a curve of "competition" of binding by an allosteric modulator, based on the ternary complex model. Note that this model assumes the allosteric modulator is present in excess, so is not depleted by binding to the receptors. Since it binds to a different site than the radioligand, the term 'competition' is not apt, but we list it here because the experimental design is the same as used for competitive binding.

**Step by step**

Create an XY data table. Enter the logarithm of the concentration of the unlabeled modulator (in nM) into X, and specific binding into Y in any convenient units.

From the data table, click Analyze, choose nonlinear regression, choose the panel of Competition Binding equations, and choose *Allosteric modulator titration*.

You must constrain two parameters to constant values based on your experimental design:

- RadioligandNM is the concentration of labeled ligand in nM. A single concentration of radioligand is used for the entire experiment.
- HotKdNM is the equilibrium dissociation constant of the labeled ligand in nM.

Also consider constraining Y0 (radioligand binding in the absence of modulator) to a constant value.

**Model**

\[
\begin{align*}
\text{AlloNM} &= 10^{(X+9)} \\
\text{KbNM} &= 10^{(\text{logKb} + 9)} \\
\alpha &= 10^{\text{logAlpha}} \\
\text{KAppNM} &= \text{HotKDnm} * ((1 + (\text{AlloNM}/\text{KBNM})) / (1 + \alpha * (\text{AlloNM}/\text{KBNM}))) \\
\text{HotOccupancy} &= \text{RadioligandNM} / (\text{RadioligandNM} + \text{HotKDnm}) \\
Y &= (\text{Y0}/\text{HotOccupancy}) * (\text{RadioligandNM} / (\text{RadioligandNM} + \text{KAppNM}))
\end{align*}
\]
Interpret the parameters

**\(K_b\)** is the equilibrium dissociation constant (Molar) of modulator binding.

**Alpha** is the ternary complex constant. When alpha=1.0, the modulator does not alter binding. If alpha is less than 1.0, then the modulator reduces ligand binding. If alpha is greater than 1.0, then the modulator increases binding.

**\(Y_0\)** is the radioligand binding in the absence of modulator. Consider constraining this to a constant value.

Notes

This model is designed to analyze data when the unlabeled compound works via an allosteric site. Since the labeled and unlabeled ligands are acting via different sites, it is inappropriate (and incorrect) to refer to these types of experiments as “competition binding assays”. In some cases, in fact, the allosteric modulator enhances radioligand binding.

The model is written to fit the logarithm of alpha, rather than alpha itself. This is because alpha is asymmetrically (all values from 0 to 1 mean that the modulator decreases binding, while all values from 1 to infinity mean that the modulator enhances binding. On a log scale, its values are more symmetrical, so the confidence interval computed on a log scale (as Prism does) are more accurate.
The Y axis plots specific binding. Even at very high concentrations of inhibitor, the specific binding does not descend to zero. This is the nature of allosteric inhibition. If alpha is very high, then the binding is inhibited almost to zero. If alpha is not so high, then the maximum inhibition is more modest. For example, if alpha=3, the maximum inhibition is down to 33%.

This model assumes that the allosteric modulator is present in excess, so the concentration you added is very close to its free concentration. This model does not apply when the concentration of allosteric modulator is limiting (as it is when G proteins alter agonist binding to many receptors). No explicit model can handle this situation. You need to define the model with an implicit equation (Y on both sides of the equals sign) and Prism cannot handle such equations.

Reference


3.6.8 Receptor binding - Kinetics

Kinetics refers to changes over time. Kinetic binding experiments are used to determine the association and dissociation rate constants.

3.6.8.1 Key concepts: Kinetics of binding

Law of mass action

Kinetic binding experiments are used to determine the association and dissociation rate constants. All the analyses assume that binding follows the law of mass action:

\[
\text{Receptor} + \text{Ligand} \xrightleftharpoons[K_{\text{off}}]{K_{\text{on}}} \text{Receptor} \cdot \text{Ligand}
\]

At any given time, the rate at which receptor-ligand complexes form is proportional to the radioligand concentration and the number of
receptors still unoccupied. The rate of dissociation is proportional to the concentration of receptor-ligand complexes.

**Rate of dissociation**

A dissociation binding experiment measures the “off rate” for radioligand dissociating from the receptor. Initially ligand and receptor are allowed to bind, perhaps to equilibrium. At that point, you need to block further binding of radioligand to receptor (by adding an unlabeled drug or by dilution) so you can measure the rate of dissociation, which follows a one-phase exponential decay with a rate constant equal to the rate of radioligand dissociation.

**Rate of association**

In an association experiment, you add radioligand and measure specific binding at various times thereafter.

Binding increases over time until it plateaus. This plateau is not the same as the Bmax. The plateau in an association experiment depends on the concentration of radioligand used, while the Bmax is extrapolated to an infinite concentration of radioligand.

The rate at which binding increases is determined by three factors (as well as experimental conditions such as pH and temperature):

- The association rate constant, $k_{on}$ or $k+1$. This is what you are trying to determine.

- The concentration of radioligand. If you use more radioligand, the system equilibrates faster.

- The dissociation rate constant, $k_{off}$ or $k-1$. Some people are surprised to see that the observed rate of association depends in part on the dissociation rate constant. During the incubation, radioligand both binds to and dissociates from receptors. The system reaches equilibrium when the two rates are equal. The observed rate of association measures how long it takes to reach equilibrium. If the radioligand dissociates quickly from the receptor, equilibrium will be reached faster (but with less binding).

It is not possible, therefore, to fit a simple association experiment to determine the association rate constant unless you constrain the value of
the dissociation rate constant, measure association kinetics with two or more concentrations of radioligand, or determine the rate of association and dissociation in one experiment.

3.6.8.2 Equation: Dissociation kinetics

Introduction

A dissociation binding experiment measures the “off rate” for radioligand dissociating from the receptor. Initially ligand and receptor are allowed to bind, perhaps to equilibrium. At that point, you need to block further binding of radioligand to receptor so you can measure the rate of dissociation. There are several ways to do this:

- If the tissue is attached to a surface, you can remove the buffer containing radioligand and replace with fresh buffer without radioligand.
- Spin the suspension and resuspend in fresh buffer.
- Add a very high concentration of an unlabeled ligand. If this concentration is high enough, it will instantly bind to nearly all the unoccupied receptors and thus block binding of the radioligand.
- Dilute the incubation by a large factor, at least 100 fold dilution. This will reduce the concentration of radioligand by that factor. At such a low concentration, new binding of radioligand will be negligible. This method is only practical when you use a fairly low concentration of radioligand so its concentration after dilution is far below its Kd for binding.

You then measure binding at various times after that to determine how rapidly the radioligand falls off the receptors.

Step by step

Create an XY data table. Enter time into X, and total binding into Y. If you have several experimental conditions, place the first into column A, the second into column B, etc.

From the table of specific binding, click Analyze, choose nonlinear regression, choose the panel of Kinetics Binding equations, and choose Dissociation - One phase exponential decay.
Model

\[ Y = (Y_0 - NS) \cdot \exp(-K \cdot X) + NS \]

Interpret the parameters

**Y₀** is the binding at time zero, in the units of the Y axis.

**NS** is the binding (nonspecific) at infinite times, in the units of the Y axis.

**K** is the rate constant in inverse units of the X axis. The half-life equals the ln(2) divided by K.

Checking for cooperativity

If the law of mass action applies, binding of a ligand to one binding site does not alter the affinity of another binding site. This also means that dissociation of a ligand from one site should not change the dissociation of ligand from other sites. To test this assumption, compare the dissociation rate after initiating dissociation by infinite dilution with the dissociation rate when initiated by addition of a large concentration of unlabeled drug. If the radioligand is bound to multiple noninteracting binding sites, the dissociation will be identical in both experimental protocols as shown in the left figure below. Note that the Y axis is shown using a log scale. If there were a single binding site, you would expect the dissociation data to appear linear on this graph. With two binding sites, the graph is curved even on a log axis.
The right figure shows ideal dissociation data when radioligand is bound to interacting binding sites with negative cooperativity. The data are different depending on how dissociation was initiated. If dissociation is initiated by infinite dilution, the dissociation rate will change over time. The dissociation of some radioligand will leave the remaining ligand bound more tightly. When dissociation is initiated by addition of cold drug, all the receptors are always occupied by ligand (some hot, some cold) and dissociation occurs at its maximal unchanging rate.

3.6.8.3 Equation: Association kinetics (one ligand concentration)

Introduction

When you measure the association rate of a radioligand, the rate at which the binding equilibrates depends not only on the association rate constant and the amount of ligand you used, but also on its dissociation rate constant. The only way to fit the association rate constant by analyzing association data from one concentration of radioligand, is to constrain the dissociation rate constant to a value you determined in a different experiment.

Alternative methods of determining an association rate constant are to globally fit data obtained with multiple radioligand concentrations, or to analyze an experiment that measures both association and dissociation rate sequentially.

Step by step

Create an XY data table. Enter time in minutes into X, and specific binding into Y.

From the table of specific binding, click Analyze, choose nonlinear regression, choose the panel of Kinetics Binding equations, and choose Association kinetics - One conc. of hot.

You must constrain Hotnm ([radioligand] in nM) and Koff (dissociation rate constant, in inverse minutes) to constant values.
Model

\[K_d = \frac{K_{off}}{K_{on}}\]
\[L = Ho \times nm \times 1e-9\]
\[K_{ob} = K_{on} \times L + K_{off}\]
\[\text{Occupancy} = \frac{L}{(L + K_d)}\]
\[Y_{max} = \text{Occupancy} \times B_{max}\]
\[Y = Y_{max} \times (1 - \exp(-1 \times k_{ob} \times X))\]

Interpret the parameters

**Kon** is the association rate constant, in units of M⁻¹ min⁻¹

**Kd** is the equilibrium binding constant, in Molar, computed as \(K_{off}/K_{on}\)

**Bmax** is the maximal binding at equilibrium, extrapolated to maximal radioligand concentration, in the units used to enter the Y values.

Notes

3.6.8.4 Equation: Association kinetics (two ligand concentrations)

Introduction

You cannot determine \(k_{on}\) from an association binding measured at a single concentration of radioligand. The observed association rate depends on the association rate constant, the amount of ligand you used, and its dissociation rate constant. With one concentration of radioligand, the results are ambiguous.

If you perform an association kinetic experiments with multiple radioligand concentration, you can globally fit the data to the association kinetic model to derive a single best-fit estimate for \(k_{on}\) and one for \(k_{off}\).
Shown below is an example of an association kinetic experiment conducted using two concentrations of radioligand. All other conditions (temperature, pH, etc.) were the same for both runs, of course. Times were entered into the X column, specific binding for one concentration of radioligand were entered into the first (A) Y column, and binding for the other concentration were entered into column B.

**Step by step**

Create an XY data table. Enter time in minutes into X, and total binding into Y. Enter binding at one concentration of radioligand into the column A, binding at another concentration into column B, etc. Enter the concentrations, in nM, into the column titles.

From the table of specific binding, click Analyze, choose nonlinear regression, choose the panel of Kinetics Binding equations, and choose *Association - Two or more conc. of hot*.

**Model**

\[
\begin{align*}
K_d &= \frac{K_{off}}{K_{on}} \\
L &= Hotnm \times 1e-9 \\
K_{ob} &= K_{on}L + K_{off} \\
Occupancy &= \frac{L}{(L + K_d)} \\
Y_{max} &= Occupancy \times B_{max} \\
Y &= Y_{max} \times (1 - \exp(-1 \times K_{ob} \times X)) 
\end{align*}
\]

**Interpret the parameters**

*Koff* is the dissociation rate constant in min\(^{-1}\).

*Kon* is the association rate constant, in units of M\(^{-1}\) min\(^{-1}\).
**Kd** is the equilibrium binding constant, in Molar, computed as $K_{off}/K_{on}$

**Bmax** is the maximal binding at equilibrium, at maximal radioligand concentration, in the units you used to enter $Y$.

**Notes**

According to the law of mass action, the ratio of $k_{off}$ to $k_{on}$ is the Kd of receptor binding:

$$K_d = \frac{k_{off}}{k_{on}}$$

Compare the Kd calculated this way (from kinetic experiments) with the Kd determined from a saturation binding curve. If binding follows the law of mass action, the two Kd values should be indistinguishable.

### 3.6.8.5 Equation: Association then dissociation

**Introduction**

You cannot determine the association rate constant by simply observing the association of a single concentration of radioligand. The rate at which a ligand reaches equilibrium is determined not only by the association rate constant and the ligand concentration, but also by the dissociation constant.

One way to determine the association rate constant is to globally fit data obtained with two different concentrations of radioligand. An alternative approach, explained here is to measure association and dissociation in one experiment.

Add a radioligand and measure total binding at multiple time points, then at Time0 initiate dissociation (by adding an antagonist or by massive dilution) and then measure dissociation at various times.

**Step by step**

Create an XY data table. Enter time in minutes into X, and total binding into Y.
From the table of specific binding, click Analyze, choose nonlinear regression, choose the panel of Kinetics Binding equations, and choose *Association then dissociation*.

Constrain HotNM (\([\text{radioigand in nM}]\) and Time0 (time at which dissociation was initiated) to constant values. If you entered specific binding into the Y column, also constrain NS to a constant value of zero.

**Model**

\[
\begin{align*}
\text{Radioligand} &= \text{HotNM} \times 1e-9 \\
K_{ob} &= [\text{Radioligand}] \times K_{on} + K_{off} \\
K_{d} &= K_{off} / K_{on} \\
Eq &= \text{Bmax} \times \text{radioligand} / (\text{radioligand} + K_{d}) \\
\text{Association} &= Eq \times (1 - \exp(-1 \times K_{ob} \times X)) \\
Y_{at Time0} &= Eq \times (1 - \exp(-1 \times K_{ob} \times Time0)) \\
\text{Dissociation} &= Y_{at Time0} \times \exp(-1 \times K_{off} \times (X - Time0)) \\
Y &= \text{IF}(X < Time0, \text{Association}, \text{Dissociation}) + NS
\end{align*}
\]

**Interpret the parameters**

\(K_{off}\) is the dissociation constant in \(\text{min}^{-1}\).

\(K_{on}\) is the association constant in inverse minutes multiplied by inverse concentration.

\(K_{D}\) is computed from \(K_{off}/K_{on}\). Expressed in Molar units.

\(B_{max}\) is the maximum binding at equilibrium with maximum concentration of radioligand, in units of Y axis. Unless you used a very
high concentration of radioligand, Bmax will have a wide confidence interval as the experiment is not designed to determine Bmax.

**NS** is the nonspecific binding, in units of the Y axis. It is the Y value at time 0, and also the Y value at very late times after all the ligand has dissociated.

**Notes**

3.6.8.6 Equation: Kinetics of competitive binding

**Introduction**

Kinetics experiments can determine the dissociation and association rate constants (off-rate and on-rate) of an unlabeled compound. Add labeled and unlabeled ligand together and measure the binding of the labeled ligand over time. Fit to the appropriate model described below, constraining the rate constants of the labeled ligand to constant values determined from other experiments, and fit the rate constants of the unlabeled compound.

Using only a single concentration of labeled and radioligand, it is very hard to determine the rate constants with any reasonable precision. But measure the kinetics at two (or more) concentrations of the unlabeled ligand, and the results are much more precise.

**Step by step**

Create an XY data table. Enter time in minutes into X, and specific binding in cpm into Y. Enter the binding for one concentration of of the unlabeled ligand in column A, and another concentration in column B, etc.. Enter the concentrations, in nM, into the column titles.

From the table of specific binding, click Analyze, choose nonlinear regression, choose the panel of Kinetics Binding equations, and choose *Kinetics of competitive binding*.

Constrain k1 and k2 to constant values determined from kinetic binding experiments. k1 is the association rate constant of the hot ligand in M\(^{-1}\) min\(^{-1}\) and k2 is its dissociation rate constant in units of min\(^{-1}\).

Also constrain L to be a constant value equal to the concentration of labeled ligand in nM.
I is constrained to be a column constant whose value comes from the column titles.

**Model**

\[
\begin{align*}
KA &= K_1 \cdot L \cdot 10^{-9} + k_2 \\
KB &= K_3 \cdot I \cdot 10^{-9} + K_4 \\
S &= \sqrt{(KA - KB)^2 + 4 \cdot K_1 \cdot K_3 \cdot L \cdot I \cdot 10^{-18}} \\
KF &= 0.5 \cdot (KA + KB + S) \\
KS &= 0.5 \cdot (KA + KB - S) \\
DIFF &= K_F - K_S \\
Q &= B_{max} \cdot K_1 \cdot L \cdot 10^{-9} / DIFF \\
Y &= Q \cdot (K_4 \cdot DIFF / (K_F \cdot K_S) + (K_4 - K_f) / K_F) \cdot \exp(-K_F \cdot X) - (K_4 - K_S) / K_S \cdot \exp(-K_S \cdot X))
\end{align*}
\]

**Interpret the parameters**

\(k_3\) is the association rate constant of unlabeled ligand in \(\text{M}^{-1} \text{min}^{-1}\).

\(k_4\) is the dissociation rate constant of unlabeled ligand in \(\text{min}^{-1}\).

\(B_{max}\) is the total number of receptors. Either leave as a variable or set to a constant you know from other experiments. The \(B_{max}\) is the maximum binding at equilibrium with a very high concentration of radioligand. It is usually much larger than any binding seen in the experiment.

All three parameters are constrained to be shared, so Prism fits one value of each of the three parameters for all the data sets.
Notes

- This equation does not account for ligand depletion. It assumes that only a small fraction of radioligand binds to receptors, so that the free concentration of radioligand is very close to the added concentration.

- This method will only give reliable results if you have plenty of data points at early time points.

- The ratio K4/K3 is the equilibrium dissociation constant of the cold ligand in Molar. You should compare this value (determined via kinetics) with the same value determined by equilibrium competition.

Reference

This method was described by Motulsky and Mahan in Molecular Pharmacology 25:1-9, 1984.

3.6.9 Enzyme kinetics -- Key concepts
3.6.9.1 Key concepts: Terminology

What is an enzyme?

Living systems depend on chemical reactions which, on their own, would occur at extremely slow rates. Enzymes are catalysts that reduce the needed activation energy so these reactions proceed at rates that are useful to the cell. The study of enzyme kinetics can help us understand the function and regulation of enzymes.

Enzyme progress curves

In most cases, an enzyme converts one chemical (the substrate) into another (the product). A graph of product concentration vs. time follows three phases marked on the graph below.
1. At very early time points (usually less than a second), the rate of product accumulation increases over time. Special techniques, not available in Prism, are needed to study the early kinetics of enzyme action. The graph above exaggerates this first phase.

2. For an extended period of time, the product concentration increases linearly with time. All the analyses built-in to Prism use data collected during this second phase.

3. At later times, the substrate is depleted, so the curve starts to level off. Eventually the concentration of product reaches a plateau.

It is very difficult to fit a curve to these kind of data. The model simply cannot be reduced to an equation that expresses product concentration as a function of time. To fit these kind of data (called an enzyme progress curve) you need to use a program that can fit data to a model defined by differential equations or by an implicit equation. For more details, see RG Duggleby, *Analysis of Enzyme Reaction Progress Curves by Nonlinear Regression*, Methods in Enzymology, 249: 61-60, 1995.

Rather than fit the enzyme progress curve, most analyses of enzyme kinetics (including all those built-in to Prism) measure product at a single time point. Analyses assume that the time point you chose is on the linear (second) phase of product accumulation and ignore the nonlinear first phase (which is usually very short). Therefore, if you divide the amount of product produced by the time the reaction was allowed to proceed, you compute the amount of product formed per unit time, which is the enzyme velocity.
Terminology

The terminology can be confusing. Note these confusing points:

- As mentioned above, almost all studies of enzyme "kinetics" are done by collecting data at a single time point. The X axis is substrate (or inhibitor) concentration, not time.

- The second phase shown in the graph above is often called the "initial rate", a phrase that makes sense only if you ignore the short transient phase that preceded it.

- That second phase is also called "steady state", because the concentration of enzyme-substrate complex doesn't change during that phase. However, the concentration of product accumulates, so the system is not truly at steady state until, much later, the concentration of product truly doesn't change over time.

3.6.9.2 Key concepts: Assumptions

Standard analyses of enzyme kinetics (the only kind discussed here) assume:

- The production of product is linear with time during the time interval used.

- The concentration of substrate vastly exceeds the concentration of enzyme. This means that the free concentration of substrate is very close to the concentration you added, and that substrate concentration is essentially constant throughout the assay.

- A single enzyme forms the product.

- There is negligible spontaneous creation of product without enzyme.

With the exception of the built-in allosteric model, Prism's built-in equations also assume:

- No cooperativity. Binding of substrate to one enzyme binding site doesn't influence the affinity or activity of an adjacent site.
Neither substrate nor product acts as an allosteric modulator to alter the enzyme velocity.

### 3.6.10 Enzyme kinetics - Substrate vs. velocity

#### 3.6.10.1 Key concepts: Substrate vs. velocity

**Fitting a substrate-velocity curve**

The simplest experiment in enzyme kinetics is to vary the substrate concentration and measure enzyme velocity (also called enzyme activity).

The standard way to fit these data is to fit the Michaelis-Menten model to determine the Vmax (maximum enzyme velocity) and its Km (the concentration of substrate needed to get half-maximal velocity).

The Vmax equals the product of the concentration of active enzyme sites times the turnover rate, kcat. This is the number of substrate molecules each enzyme site can convert to product per unit time. If you know the concentration of enzyme, you can fit the curve to determine kcat and Km. The curve will be identical to the Michaelis-Menten fit.

If the enzyme has cooperative subunits, the graph of enzyme velocity as a function of substrate concentration will appear sigmoidal. Prism offers one empirical equation for fitting sigmoidal substrate-velocity curves.

#### 3.6.10.2 Equation: Michaelis-Menten model

**Introduction**

The most common kind of enzyme kinetics experiment is to vary the concentration of substrate and measure enzyme velocity. The goal is to determine the enzyme's Km (substrate concentration that yield a half-maximal velocity) and Vmax (maximum velocity). If your goal is to determine the turnover number kcat, rather than the Vmax, use an alternative version of the equation.
**Step by step**

Create an XY data table. Enter substrate concentration into X, and enzyme velocity into Y. If you have several experimental conditions, place the first into column A, the second into column B, etc.

You can also choose Prism's sample data: Enzyme kinetics -- Michaelis-Menten.

After entering data, click Analyze, choose nonlinear regression, choose the panel of enzyme kinetics equations, and choose *Michaelis-Menten enzyme kinetics*.

**Model**

\[ Y = \frac{V_{\text{max}} \times X}{(K_m + X)} \]

**Interpret the parameters**

- **Vmax** is the maximum enzyme velocity in the same units as Y. It is the velocity of the enzyme extrapolated to very high concentrations of substrate, so its value is almost always higher than any velocity measured in your experiment.
**Km** is the Michaelis-Menten constant, in the same units as X. It is the substrate concentration needed to achieve a half-maximum enzyme velocity.

**Create a Lineweaver-Burk plot**

Before nonlinear regression was available, investigators had to transform curved data into straight lines, so they could analyze with linear regression. One way to do this is with a Lineweaver-Burk plot, which plots the reciprocal of substrate concentration vs. the reciprocal of enzyme velocity.

![Lineweaver-Burk plot diagram](image)

If you create a Lineweaver-Burk plot, use it only to display your data. Don't use the slope and intercept of a linear regression line to determine values for Vmax and Km. If you do this, you won't get the most accurate values for Vmax and Km. The problem is that the transformations (reciprocals) distort the experimental error, so the double-reciprocal plot does not obey the assumptions of linear regression. Use nonlinear regression to obtain the most accurate values of Km and Vmax.

To create a Lineweaver-Burk plot with Prism, use the Transform analysis, then choose the panel of biochemistry and pharmacology transforms.

To create a Lineweaver-Burke line corresponding to the nonlinear regression fit, follow these steps:

1. Create a new XY data table, with no subcolumns.

2. Into row 1 enter $X = -1/K_m$, $Y = 0$ (previously determined by nonlinear regression).

3. Into row 2 enter $X = 1/S_{\text{min}}$ ($S_{\text{min}}$ is the smallest value of [substrate] you want to include on the graph) and $Y = (1/V_{\text{max}})(1.0 + K_m/S_{\text{min}})$. 
4. Note the name of this data table. Perhaps rename it to something appropriate.

5. Go to the Lineweaver-Burke graph.

6. Drag the new table from the navigator and drop onto the graph.

7. Double-click on one of the new symbols for that data set to bring up the Format Graph dialog.

8. Choose to plot no symbols, but to connect with a line.

Notes

- See the list of assumptions of all analyses of enzyme kinetics.

- This equation fits exactly the same curve as the equation that fits the turnover number \(K_{\text{cat}}\) rather than the \(V_{\text{max}}\). The product of \(K_{\text{cat}}\) times \(E_t\) (the concentration of enzyme sites) equals the \(V_{\text{max}}\), so if you know \(E_t\), Prism can fit \(k_{\text{cat}}\).

- This equation is a special case of the equation for allosteric enzymes. That allosteric model adds an additional parameter: the Hill slope \(h\). When \(h\) equals 1.0, the two models are identical.

- Note that \(K_m\) is not a binding constant that measures the strength of binding between the enzyme and substrate. Its value takes into account the affinity of substrate for enzyme, and also the rate at which the substrate bound to the enzyme is converted to product.

3.6.10.3 Equation: Determine \(k_{\text{cat}}\)

Introduction

\(K_{\text{cat}}\) is the turnover number -- the number of substrate molecule each enzyme site converts to product per unit time. If you know the concentration of enzyme sites, you can fit \(K_{\text{cat}}\) instead of \(V_{\text{max}}\) when analyzing a substrate vs. velocity curve.

The model

\[ Y = E_t \cdot k_{\text{cat}} \cdot X / (K_m + X) \]
\( X \) is the substrate concentration.

\( Y \) is enzyme velocity.

**\( \text{kcat} \)** is the turnover number, the number of times each enzyme site converts substrate to product per unit time. This is expressed in the inverse of the time units of the \( Y \) axis. For example, if \( Y \) is in micromoles of substrate per minute, then \( \text{kcat} \) is the number of molecules of substrate produced per catalytic site per minute.

**\( \text{Km} \)** is the Michaelis-Menten constant, in the same units as \( X \). It is the substrate concentration needed to achieve a half-maximum enzyme velocity.

**\( \text{Et} \)** is the concentration of enzyme catalytic sites. If the enzyme has multiple subunits, note that \( \text{Et} \) is the concentration of catalytic sites, which can be larger than the concentration of enzyme molecules. The \( Y \) values are entered in units of concentration per time, and \( \text{Et} \) must be entered in those same concentration units.

**\( \text{Vmax} \)** is the maximum enzyme velocity in the same units as \( Y \). It is not directly shown in the model above. It is the velocity of the enzyme extrapolated to very high concentrations of substrate, so is almost
always higher than any velocity measured in your experiment. It is computed by multiplying Et times kcat.

**Relationship to the Michaelis-Menten model**

The curve shown above is identical to the curve defined by the Michaelis-Menten model. You'll get identical curves when you fit either model to your data, and identical values for Km.

The Michaelis-Menten model finds the Vmax, which is the maximum enzyme velocity extrapolated out to very high concentrations of substrate. It is expressed in the same units you used to enter your Y values (enzyme activity). Usually it is straightforward to express this (or convert to) moles/minute/mg of protein. The Vmax is determined by how many enzyme sites are present (Et) and the rate at which the enzyme can convert substrate to product (kcat).

If you know the concentration of enzyme sites you've added to the assay (Et) then you can fit the catalytic constant Kcat using the model above.

When calculating Kcat, the concentration units cancel out, so Kcat is expressed in units of inverse time. It is the turnover number -- the number of substrate molecule each enzyme site converts to product per unit time.

**Fitting the model with Prism**

1. Create an XY data table. Enter substrate concentration into X, and enzyme velocity into Y. If you have several experimental conditions, place the first into column A, the second into column B, etc. You can also choose Prism's sample data: Enzyme kinetics -- Michaelis-Menten.

2. After entering data, click Analyze, choose nonlinear regression, choose the panel of enzyme kinetics equations, and choose \( \text{Kcat} \).

3. You must constrain \( \text{Et} \) to a constant value, based on other experiments. To constrain the value of Et, go to the Constrain tab of the nonlinear regression dialog, make sure that the drop down next to Et is set to "Constant equal to" and enter the value. For the sample data, enter 100 as the value of Et.

   If you don't know the value of Et, you cannot fit the kcat, but instead should fit the Vmax. It is not possible for Prism to fit both the kcat
and Et, as the two parameters are intertwined, and a substrate-velocity curve gives no information about their individual values.

4. With the sample data, Prism reports that $K_m = 5.886$ with a 95% confidence interval ranging from 3.933 to 7.839. The best fit value of $k_{cat}$ is 13.53 with a 95% confidence interval ranging from 11.97 to 15.09.

**Notes**

- See the list of assumptions of all analyses of enzyme kinetics.
- This equation fits exactly the same curve as the equation that fits $V_{max}$, rather than the turnover number $K_{cat}$. The product of $K_{cat}$ times $E_t$ (the concentration of enzyme sites) equals the $V_{max}$.
- This equation is related to the equation for allosteric enzymes. That allosteric model adds an additional parameter: the Hill slope $h$. When $h$ equals 1.0, the two models are identical.

### 3.6.10.4 Equation: Allosteric sigmoidal

**Introduction**

If the enzyme has cooperative subunits, the graph of enzyme velocity as a function of substrate concentration will appear sigmoidal. Prism offers one empirical equation for fitting sigmoidal substrate-velocity curves. Read advanced books on enzyme kinetics for alternative methods based on molecular models of allosteric action.

**How to enter data**

Create an XY data table. Enter substrate concentration into $X$, and enzyme velocity into $Y$. If you have several experimental conditions, place the first into column A, the second into column B, etc.

After entering data, click Analyze, choose nonlinear regression, choose the panel of enzyme kinetics equations, and choose *Allosteric sigmoidal enzyme kinetics*. 
The model

\[ Y = \frac{V_{\text{max}} \times X^h}{(K_{\text{half}}^h + X^h)} \]

Interpret the parameters

**Vmax** is the maximum enzyme velocity in the same units as \( Y \). It is the velocity of the enzyme extrapolated to very high concentrations of substrate, and therefore is almost always higher than any velocity measured in your experiment.

**Khalf** is the concentration of substrate that produces a half-maximal enzyme velocity. It is the EC50.

**h** is the Hill slope. When \( h = 1 \), this equation is identical to the standard Michaelis-Menten equation\[^{[36]}\]. When it is greater than 1.0, the curve is sigmoidal due to positive cooperativity. The variable \( h \) does not always equal the number of interacting binding sites, but its value can not exceed the number of interacting sites. Think of \( h \) as an empirical measure of the steepness of the curve and the presence of cooperativity.

**Kprime** is related to the Km. It is computed as \( K_{\text{half}}^h \), and is expressed in the same units as \( X \).

An alternative form of the equation

An alternative version of the equation (equation 5.47 in the book by Dr. Copeland referenced below) fits Kprime.

\[ Y = \frac{V_{\text{max}} \times X^h}{(K_{\prime} + X^h)} \]
Note that Kprime in this equation equals Khalf^h in the equation built into Prism. The two models generate exactly the same curve, and simply have alternative methods for reporting the parameters. Prism reports both Kprime and Khalf.

Reference


3.6.11 Enzyme kinetics -- Inhibition

Many drugs work by inhibiting enzyme activity, either by preventing the substrate from binding to the enzyme, or by stabilizing the enzyme-substrate complex so as to slow formation of product. To distinguish between the models of enzyme inhibition and determine the Ki of the inhibitor, measure substrate-velocity curves in the presence of several concentrations of inhibitor (including one curve with no inhibitor).

3.6.11.1 Key concepts: Enzyme inhibition

Many drugs work by inhibiting enzyme activity, either by preventing the substrate from binding to the enzyme, or by stabilizing the enzyme-substrate complex so as to slow formation of product. To distinguish between the models of enzyme inhibition and determine the Ki of the inhibitor, measure substrate-velocity curves in the presence of several concentrations of inhibitor (including one curve with no inhibitor).

Prism can fit your data to six models of enzyme inhibition:

- A **competitive** inhibitor reversibly binds to the same site as the substrate, so its inhibition can be entirely overcome by using a very high concentration of substrate. The maximum velocity of the enzyme doesn't change (if you give it enough substrate), but it takes more substrate to get to half maximal activity. The substrate-velocity curve is shifted to the left but not down.

- A **noncompetitive** inhibitor binds with equal affinity to the enzyme, and the enzyme-substrate complex. The inhibition is not surmountable
by increasing substrate concentration. The substrate-velocity curve is shifted down but neither to the right or left.

- An **uncompetitive** inhibitor reversibly binds to the enzyme-substrate complex, but not to the enzyme itself. This reduces both the effective Vmax and the effective Km. The substrate-velocity curve is shifted down and to the left.

- The **mixed** model is a general model that includes competitive, noncompetitive and uncompetitive models as special cases. The model has one more parameter than the others, and the extra parameter (alpha) determines the degree to which the binding of inhibitor changes the affinity of the enzyme for substrate.

- **Substrate inhibition**. In some cases, the substrate of an enzyme also inhibits the enzyme by binding to a second site on the enzyme.

- **Tight inhibition**. This model accounts for tight binding, so it does not assume that the free concentration of inhibitor equals the total concentration.

Reference


### 3.6.11.2 Equation: Competitive inhibition

#### Introduction

A competitive inhibitor reversibly binds to the same site as the substrate, so its inhibition can be entirely overcome by using a very high concentration of substrate. The Vmax doesn't change, and the effective Km increases. You can determine the Ki of a competitive inhibitor by measuring substrate-velocity curves in the presence of several concentrations of inhibitor.

#### Step by step

Create an XY data table. Enter substrate concentration into the X column, and enzyme activity into the Y columns. Each data set (Y column) represents data collected in the presence of a different concentration of inhibitor, starting at zero. Enter these concentrations
into the column titles. Be sure to enter concentrations, not logarithms of concentration.

Alternatively, choose the competitive enzyme inhibition sample data set.

After entering data, click Analyze, choose nonlinear regression, choose the panel of enzyme kinetics equations, and choose *Competitive enzyme inhibition*.

**The model**

\[ \text{Km}_{\text{obs}} = \text{Km} \times (1 + \frac{[I]}{K_i}) \]

\[ Y = \frac{V_{\text{max}} \times X}{(\text{Km}_{\text{obs}} + X)} \]

The constant \( I \) is the concentration of inhibitor, a value you enter into each column title. This is constrained to equal a data set constant.

The parameters \( V_{\text{max}}, \text{Km} \) and \( K_i \) are shared, so Prism fits one best-fit value for the entire set of data.

![Graph](image-url)

**Interpreting the parameters**

- \( K_i \) is the inhibition constant, expressed in the same units as \( I \), which you entered into the column titles.

- \( V_{\text{max}} \) is the maximum enzyme velocity, in the absence of inhibitor, expressed in the same units as \( Y \).
**Km** is the Michaelis-Menten constant, expressed in the same units as X. It describes the interaction of substrate and enzyme in the absence of inhibitor.

If the data don't fit the model well, consider instead fitting to a noncompetitive or uncompetitive model. Or fit to the more general equation for mixed-model inhibition.

**Reference**


**3.6.11.3 Equation: Noncompetitive inhibition**

**Introduction**

A noncompetitive inhibitor reversibly binds to both the enzyme-substrate complex, and the enzyme itself. This means that the effective Vmax decreases with inhibition but the Km does not change. You can determine the Ki of a competitive inhibitor by measuring substrate-velocity curves in the presence of several concentrations of inhibitor.

The term 'noncompetitive' is used inconsistently. It is usually used as defined above, when the inhibitor binds with identical affinity to the free enzyme and the enzyme-substrate complex. Sometimes, however, the term 'noncompetitive' is used more generally, when the two binding affinities differ, which is more often called mixed-model inhibition.

**Step by step**

Create an XY data table. Enter substrate concentration into the X column, and enzyme activity into the Y columns. Each data set (Y column) represents data collected in the presence of a different concentration of inhibitor, starting at zero. Enter these concentrations into the column titles. Be sure to enter concentrations, not logarithms of concentration.

After entering data, click Analyze, choose nonlinear regression, choose the panel of enzyme kinetics equations, and choose Noncompetitive enzyme inhibition.
The model

\[ V_{\text{max inh}} = \frac{V_{\text{max}}}{1 + \frac{I}{K_i}} \]
\[ Y = \frac{V_{\text{max inh}} \times X}{K_m + X} \]

The constant \( I \) is the concentration of inhibitor, a value you enter into each column title. This is constrained to be a data set constant.

The parameters \( \textbf{Vmax}, \textbf{Km} \) and \( \textbf{Ki} \) are shared, so Prism fits one best-fit value for the entire set of data.

Interpreting the parameters

\( \textbf{Vmax} \) is the maximum enzyme velocity without inhibitor, expressed in the same units as \( Y \).

\( \textbf{Km} \) is the Michaelis-Menten constant (without inhibitor), expressed in the same units as \( X \). It describes the interaction of substrate and enzyme in the absence of inhibitor.

\( \textbf{Ki} \) is the inhibition constant, expressed in the same units as \( I \), which you entered.

If the data don't fit the model well, consider instead fitting to a competitive or uncompetitive model. Or fit to the more general equation for mixed-model inhibition.
3.6.11.4 Equation: Uncompetitive inhibition

Introduction

An uncompetitive inhibitor binds to the enzyme-substrate complex, but not the free enzyme. This reduces both the effective Vmax and the effective Km. The substrate-velocity curve is shifted down and to the left.

You can determine the Ki of a competitive inhibitor by measuring substrate-velocity curves in the presence of several concentrations of inhibitor.

Step by step

Create an XY data table. Enter substrate concentration into the X column, and enzyme activity into the Y columns. Each data set (Y column) represents data collected in the presence of a different concentration of inhibitor, starting at zero. Enter these concentrations into the column titles. Be sure to enter concentrations, not logarithms of concentration.

After entering data, click Analyze, choose nonlinear regression, choose the panel of enzyme kinetics equations, and choose ___Uncompetitive enzyme inhibition__.

The model

\[
\begin{align*}
V_{\text{max,app}} &= \frac{V_{\text{max}}}{1 + I/\alpha K_i} \\
K_{\text{m,app}} &= \frac{K_m}{1 + I/\alpha K_i} \\
Y &= \frac{V_{\text{max,app}} X}{K_{\text{m,app}} + X}
\end{align*}
\]

The constant \( I \) is the concentration of inhibitor, a value you enter into each column title. This is constrained to equal a data set constant.

The parameters \( V_{\text{max}}, K_m \) and \( K_i \) are shared, so Prism fits one best-fit value for the entire set of data.
Interpreting the parameters

**Vmax** is the maximum enzyme velocity without inhibitor, expressed in the same units as Y.

**Km** is the Michaelis-Menten constant (without inhibitor), expressed in the same units as X. It describes the interaction of substrate and enzyme in the absence of inhibitor.

**AlphaKi** is the inhibition constant, expressed in the same units as I, which you entered into the column titles. It is the product of Ki (which is very high, because uncompetitive inhibitors don't bind to the enzyme) and alpha (which is very low). It is not possible to fit alpha and Ki separately, but only to determine their product. Some books call this product Ki'.

If the data don't fit the model well, consider instead fitting to a competitive or noncompetitive model. Or fit to the more general equation for mixed-model inhibition.

Reference

3.6.11.5 Equation: Mixed-model inhibition

**Introduction**

The mixed model is a general equation that includes competitive, uncompetitive and noncompetitive inhibition as special cases. The model has one more parameter than the others, and this parameter tells you about the mechanism of inhibition.

**Step by step**

Create an XY data table. Enter substrate concentration into the X column, and enzyme activity into the Y columns. Each data set (Y column) represents data collected in the presence of a different concentration of inhibitor, starting at zero. Enter these concentrations into the column titles. Be sure to enter concentrations, not logarithms of concentration.

After entering data, click Analyze, choose nonlinear regression, choose the panel of enzyme kinetics equations, and choose *Mixed model enzyme inhibition*.

**Model**

\[
\begin{align*}
V_{\text{max App}} &= V_{\text{max}}/(1 + I/(\text{Alpha} \times K_i)) \\
K_{\text{m App}} &= K_m 	imes (1 + I/K_i)/(1 + I/(\text{Alpha} \times K_i)) \\
Y &= V_{\text{max App}} \times X/(K_{\text{m App}} + X)
\end{align*}
\]

The parameter $I$ is the concentration of inhibitor, a value you enter into each column title. This is constrained to equal a data set constant.

The parameters *Alpha, Vmax, Km* and $K_i$ are shared, so Prism fits one best-fit value for the entire set of data.

**Interpreting the parameters**

*Vmax* is the maximum enzyme velocity without inhibitor, expressed in the same units as $Y$.

*Km* is the Michaelis-Menten constant, expressed in the same units as $X$. It describes the interaction of substrate and enzyme in the absence of inhibitor.
**Ki** is the inhibition constant, expressed in the same units as I, which you entered into the column titles.

**Alpha** determines mechanism. Its value determines the degree to which the binding of inhibitor changes the affinity of the enzyme for substrate. Its value is always greater than zero. When Alpha=1, the inhibitor does not alter binding of substrate to the enzyme, and the mixed-model is identical to noncompetitive inhibition. When alpha is very large, binding of inhibitor prevents binding of the substrate and the mixed-model becomes identical to competitive inhibition. When Alpha is very small (but greater than zero), binding of the inhibitor enhances substrate binding to the enzyme, and the mixed model becomes nearly identical to an uncompetitive model.

**Reference**


### 3.6.11.6 Equation: Substrate inhibition

**Introduction**

At high concentrations, some substrates also inhibit the enzyme activity. Substrate inhibition occurs with about 20% of all known enzymes. It happens when two molecules of substrate can bind to the enzyme, and thus block activity.

**Step by step**

Create an XY data table. Enter substrate concentration into the X column, and enzyme activity into the Y columns. If you have several experimental conditions, place the first into column A, the second into column B, etc.

After entering data, click Analyze, choose nonlinear regression, choose the panel of enzyme kinetics equations, and choose *Substrate inhibition*.

**Model**

\[ Y = \frac{V_{\text{max}} \times X}{K_m + X \times (1 + X / K_i)} \]
Parameters

**Vmax** is the maximum enzyme velocity, if the substrate didn't also inhibit enzyme activity, expressed in the same units as Y.

**Km** is the Michaelis-Menten constant, expressed in the same units as X. It describes the interaction of substrate and enzyme in the absence of inhibitor.

**Ki** is the dissociation constant for substrate binding in such a way that two substrates can bind to an enzyme. It is expressed in the same units as X.

Note: Prism 7 uses a different (improved) set of rules for initial values than did Prism 6.

Reference


3.6.11.7 Equation: Tight inhibition (Morrison equation)

**Introduction**

This equation accounts for tight binding, so it does not assume that the free concentration of inhibitor equals the total concentration.
Step by step

Create an XY data table. Enter inhibitor concentration into the X column (usually in micromolar, but any concentration units is fine), and enzyme activity into the Y columns (any units). If you have several experimental conditions, place the first into column A, the second into column B, etc.

After entering data, click Analyze, choose nonlinear regression, choose the panel of enzyme kinetics equations, and choose Morrison Ki.

Constrain Et, S and Km to constant values

You must constrain three parameters to constant values. To constrain the values, go to the Constrain tab of the nonlinear regression dialog, make sure that the drop down next to Et, S and Km is set to "Constant equal to" and enter the values.

- **Et** is the concentration of enzyme catalytic sites in the same units as the X values. If the enzyme has multiple subunits, note that Et is the concentration of catalytic sites, which can be larger than the concentration of enzyme molecules.

- **S** is the concentration of substrate you chose to use. Use the same units as the X values.

- **Km** is the Michaelis-Menten constant, expressed in the same units as X, determined in an experiment without competitor.

Prism cannot fit any of these parameters from the graph of activity vs inhibitor concentration. You must know S from your experimental design, determine Km and Et in other experiments, and constrain all three to constant values.

Model

\[ Q = (K_i \times (1 + (S/K_m))) \]
\[ Y = V_0 \times (1 - (((Et+X+Q) - (((Et+X+Q)^2 - 4*Et*X)^0.5)) / (2*Et))) \]
Interpreting parameters

\( V_0 \) is the enzyme velocity with no inhibitor, expressed in the same units as \( Y \). This is not the same as \( V_{\text{max}} \), which would require a maximal concentration of substrate.

\( K_i \) is the inhibition constant, expressed in the same units as \( X \).

Reference


3.6.11.8 Comparing models of enzyme inhibition

Comparing the fits of the different enzyme inhibition models requires thinking both about enzyme mechanism and about statistics. Some notes:

- The *mixed model* generalizes competitive, noncompetitive and uncompetitive. Each of those three models are special cases of the mixed-model.

- Comparing \( R^2 \) is an easy way to compare fits, but not a very good method.
Comparing the adjusted $R^2$ is a better way to compare fits. To do this, you need to ask for reporting of adjusted $R^2$ in the parameters dialog. But this is not a rigorous way to compare models.

Prism can compare models using two rigorous approaches, the extra sum-of-squares F test and the AICc approach.

### 3.6.12 Exponential

#### 3.6.12.1 Key concepts: Exponential equations

**What is exponential?**

Processes follow exponential models when the rate at which something is happening depends on the amount that is present.

**Exponential dissociation vs. association vs. growth**

The dissociation model always heads downhill gradually approaching a plateau.

The association model always heads uphill, and also approaches a plateau.

The growth model goes up and up and up, getting steeper, never reaching a plateau.

**Rate constants vs. time constants vs. half-lives**

In all the exponential models, one (or more in some cases) parameter describes how rapidly the process occurs. It can be expressed as a rate
constant (in units of inverse time) or as a time constant (in units of
time), or as a half-life, also in units of time.

The rate constant and time constants are simply reciprocals of each
other. Prism always fits the rate constant (k), but computes the time
constant (tau) as well and reports the standard error and confidence
interval of the time constant just as if the model had been written to fit
that constant.

The half-life equals ln(2)/k where ln is the abbreviation for natural
logarithm.

3.6.12.2 Key concepts: Derivation of exponential decay

Define X to be time, and Y to be the outcome you are measuring. Three
examples:

- Number of ligand-receptor complexes still present -- the ligand has
  not yet dissociated.
- The number of radioactive atoms that have not yet decayed.
- The concentration of drug in the plasma, not yet metabolized.

At any given time X. The rate of change of Y is proportional to Y.
Expressed as a differential equation:

\[ \frac{\Delta Y}{\Delta X} = \frac{dY}{dX} = -k \cdot Y \]

Shown as a graph:
When you integrate both sides of the equation, you get the equation for exponential decay:

\[ Y = Y_0 \exp(-kX) \]

The function \( \exp() \) takes the constant \( e \) \( (2.718...) \) to the power contained inside the parentheses.

### 3.6.12.3 Equation: One phase decay

#### Introduction

An exponential decay equation models many chemical and biological processes. It is used whenever the rate at which something happens is proportional to the amount which is left. Here are three examples:

- When ligands dissociate from receptors, the number of molecules that dissociate in any short time interval is proportional to the number that were bound at the beginning of that interval. Equivalently, each individual molecule of ligand bound to a receptor has a certain probability of dissociating from the receptor in any small time interval. That probability does not get higher as the ligand stays on the receptor longer.

- When radioactive isotopes decay, the number of atoms that decay in any short interval is proportional to the number of undecayed atoms that were present at the beginning of the interval. This means that each individual atom has a certain probability of decaying in a small time interval, and that probability is constant. The probability that any particular atom will decay does not change over time. The total decay of the sample decreases with time because there are fewer and fewer undecayed atoms.
When drugs are metabolized by the liver or excreted by the kidney, the rate of metabolism or excretion is often proportional to the concentration of drug in the blood plasma. Each drug molecule has a certain probability of being metabolized or secreted in a small time interval. As the drug concentration goes down, the rate of its metabolism or excretion goes down as well.

**Entering data**

Create an XY data table. Enter time into X, and response (binding, concentration ..) into Y. If you have several experimental conditions, place the first into column A, the second into column B, etc.

You can also choose a sample data set for exponential decay.

After entering data, click Analyze, choose nonlinear regression, choose the panel of exponential equations, and choose *One phase decay*.

**Consider constraining Plateau to a constant value of zero**

If you have subtracted off any background signal, then you know the curve has to plateau at Y=0. In this case, you should constrain the parameter Plateau to be a constant value equal to zero. To do this, go to the Constrain tab of the nonlinear regression dialog, set the drop down next to Plateau to "Constant equal to" and enter the value 0.0.

**Model**

\[ Y = (Y_0 - \text{Plateau}) \cdot e^{(-K \cdot X)} + \text{Plateau} \]
**YO** is the Y value when X (time) is zero. It is expressed in the same units as Y.

**Plateau** is the Y value at infinite times, expressed in the same units as Y.

**K** is the rate constant, expressed in reciprocal of the X axis time units. If X is in minutes, then K is expressed in inverse minutes.

**Tau** is the time constant, expressed in the same units as the X axis. It is computed as the reciprocal of K.

**Half-life** is in the time units of the X axis. It is computed as \( \ln(2)/K \).

**Span** is the difference between Y0 and Plateau, expressed in the same units as your Y values.

The derivative of an exponential decay equals \(-k\cdot Y\). So the initial rate equals \(-k\cdot Y0\).

### 3.6.12.4 Equation: Two phase decay

**Introduction**

An exponential decay equation models many chemical and biological processes. It is used whenever the rate at which something happens is proportional to the amount which is left.

A two-phase model is used when the outcome you measure is the result of the sum of a fast and slow exponential decay. This is also called a double exponential decay.

**Entering data**

Create an XY data table. Enter time into X, and response (binding, concentration ..) into Y. If you have several experimental conditions, place the first into column A, the second into column B, etc.

After entering data, click Analyze, choose nonlinear regression, choose the panel of exponential equations, and choose *Two phase decay*. 
Consider constraining Plateau to a constant value of zero

If you have subtracted off any background signal, then you know the curve has to plateau at Y=0. In this case, you should constrain the parameter Plateau to be a constant value equal to zero. To do this, go to the Constrain tab of the nonlinear regression dialog, set the drop down next to Plateau to "Constant equal to" and enter the value 0.0.

In some situations, you also know that the plateau has to be zero, even without subtracting a background. For example, in pharmacokinetics you know the drug concentration (usually) has to descend to zero at long time points.

Model

\[
\begin{align*}
\text{SpanFast} &= (Y_0 - \text{Plateau}) \times \text{PercentFast} \times 0.01 \\
\text{SpanSlow} &= (Y_0 - \text{Plateau}) \times (100 - \text{PercentFast}) \times 0.01 \\
Y &= \text{Plateau} + \text{SpanFast} \times \exp(-K_{\text{Fast}} \times X) + \text{SpanSlow} \times \exp(-K_{\text{Slow}} \times X)
\end{align*}
\]

\(Y_0\) is the Y value when X (time) is zero. It is expressed in the same units as \(Y\).

\(\text{Plateau}\) is the Y value at infinite times, expressed in the same units as \(Y\).
**Kfast** and **Kslow** are the two rate constants, expressed in reciprocal of the X axis time units. If X is in minutes, then K is expressed in inverse minutes.

**TauFast** and **TauSlow** are the two time constants, expressed in the same units as the X axis. They are computed as the reciprocals of the rate constants.

**Half-life (fast)** and **Half-life (slow)** are in the time units of the X axis. They are computed as ln(2)/K.

**PercentFast** is the fraction of the span (from Y0 to Plateau) accounted for by the faster of the two components.

**Interpretation of the parameters when you constrain Plateau to a constant value of zero**

If you have subtracted off any background signal, then you know the curve has to plateau at Y=0. You also know the curve has to descend to Y=0 in some other situations such as pharmacokinetics. In these cases, you should constrain the parameter Plateau to be a constant equal to zero.

With this constraint, it is easier to understand the model. It is the sum of two components, each defined by a rate constant and a starting Y value. The two rate constants are Kfast and Kslow. The starting Y values are defined by these equations:

\[
\begin{align*}
\text{SpanFast} &= Y0 \times \text{PercentFast} \times 0.01 \\
\text{SpanSlow} &= Y0 \times (100 - \text{PercentFast}) \times 0.01 
\end{align*}
\]

The names SpanFast and SpanSlow are probably better thought of as Y0Fast and Y0Slow.

Note the two phases are both happening at all time points. It is not correct to think that first the fast phase finishes, and only then does the slow phase start.

3.6.12.5 **Equation: Plateau followed by one phase decay**

**Introduction**
In the standard one-phase decay equation, the decay starts at time 0. This equation is used when you measure a baseline for a while, then do some experimental intervention that starts the decay at some time $X_0$.

**Entering data**

Create an XY data table. Enter time into $X$, and response (binding, concentration ..) into $Y$. If you have several experimental conditions, place the first into column A, the second into column B, etc.

After entering data, click Analyze, choose nonlinear regression, choose the panel of exponential equations, and choose *Plateau followed by one phase decay*.

**Consider constraining $X_0$ and Plateau to a constant values**

If you know the time at which you initiated the decay, you should constrain $X_0$ to that value.

If you have subtracted off any background signal, then you know the curve has to plateau at $Y=0$. In this case, you should constrain the parameter Plateau to be a constant value equal to zero.

To constrain parameters to constant values, go to the Constrain tab of the nonlinear regression dialog, set the drop down next to the parameter name to "Constant equal to" and enter the value.

**Model**

$$Y = \begin{cases} Y_0 & \text{if } X < X_0 \\ Plateau + (Y_0 - Plateau) \times \exp(-K \times (X - X_0)) & \text{otherwise} \end{cases}$$
**X0** is the time at which the decay begins. Often you will set that to a constant value based on your experimental design, but otherwise Prism can fit it. It is expressed in the same time units as X.

**Y0** is the average Y value up to time X0. It is expressed in the same units as Y.

**Plateau** is the Y value at infinite times, expressed in the same units as Y.

**K** is the rate constant, expressed in reciprocal of the X axis time units. If X is in minutes, then K is expressed in inverse minutes.

**Tau** is the time constant, expressed in the same units as the X axis. It is computed as the reciprocal of K.

**Half-life** is in the time units of the X axis. It is computed as \( \ln(2)/K \).

**Span** is the difference between Y0 and Plateau, expressed in the same units as your Y values.
3.6.12.6 Equation: Three phase decay

**Introduction**

An exponential decay equation models many chemical and biological processes. It is used whenever the rate at which something happens is proportional to the amount which is left.

A three-phase model is used when the outcome you measure is the result of the sum of a fast, medium and slow exponential decay. You need lots of data with little scatter to adequately fit a three phase model.

**Entering data**

Create an XY data table. Enter time into X, and response (binding, concentration ..) into Y. If you have several experimental conditions, place the first into column A, the second into column B, etc.

After entering data, click Analyze, choose nonlinear regression, choose the panel of exponential equations, and choose *Three phase decay*.

**Consider constraining Plateau to a constant value of zero**

If you have subtracted off any background signal, then you know the curve has to plateau at Y=0. In this case, you should constrain the parameter Plateau to be a constant value equal to zero. To do this, go to the Constrain tab of the nonlinear regression dialog, set the drop down next to Plateau to "Constant equal to" and enter the value 0.0.

**Model**

\[
Y_{Fast} = (Y_0 - \text{Plateau}) \times \text{PercentFast} \times 0.01 \times \exp(-K_{Fast} \times X) \\
Y_{Slow} = (Y_0 - \text{Plateau}) \times \text{PercentSlow} \times 0.01 \times \exp(-K_{Slow} \times X) \\
Y_{Medium} = (Y_0 - \text{Plateau}) \times (100 - \text{PercentFast} - \text{PercentSlow}) \times 0.01 \times \exp(-K_{medium} \times X) \\
Y = \text{Plateau} + Y_{Fast} + Y_{Medium} + Y_{Slow}
\]

*Y₀* is the Y value when X (time) is zero. It is expressed in the same units as Y,

**Plateau** is the Y value at infinite times, expressed in the same units as Y.
**Kfast, Kmedium and Kslow** are the rate constants, expressed in reciprocal of the X axis time units. If X is in minutes, the rate constants are expressed in inverse minutes.

**Half-life (fast, medium and slow)** are in the time units of the X axis. They are computed as \( \ln(2) \) divided by the corresponding rate constant.

**PercentFast** is the percentage of the span (from \( Y_0 \) to Plateau) accounted for by the fastest of the three components.

**PercentSlow** is the percentage of the span (from \( Y_0 \) to Plateau) accounted for by the slowest of the three components.

### 3.6.12.7 Equation: One phase association

**Introduction**

This equation describes the pseudo-first order association kinetics of the interaction between a ligand and its receptor, or a substrate and an enzyme. During each time interval a certain fraction of the unoccupied receptors become occupied. But as time advances, fewer receptors are unoccupied so fewer ligand bind and the curve levels off.

**Entering data**

Create an XY data table. Enter time into X, and response (binding, concentration ..) into Y. If you have several experimental conditions, place the first into column A, the second into column B, etc.

After entering data, click Analyze, choose nonlinear regression, choose the panel of exponential equations, and choose *One phase association*.

**Consider constraining Plateau to a constant value of zero**

If you have subtracted off any background signal, then you know the curve has to plateau at \( Y=0 \). In this case, you should constrain the parameter Plateau to be a constant value equal to zero. To do this, go to the Constrain tab of the nonlinear regression dialog, set the drop down next to Plateau to "Constant equal to" and enter the value 0.0.
Model

\[ Y = Y_0 + (\text{Plateau} - Y_0) \times (1 - \exp(-K \times x)) \]

**Y₀** is the Y value when X (time) is zero. It is expressed in the same units as Y.

**Plateau** is the Y value at infinite times, expressed in the same units as Y.

**K** is the rate constant, expressed in reciprocal of the X axis time units. If X is in minutes, then K is expressed in inverse minutes.

**Tau** is the time constant, expressed in the same units as the X axis. It is computed as the reciprocal of K.

**Half-time** is in the time units of the X axis. It is computed as \( \ln(2)/K \).

**Span** is the difference between Y₀ and Plateau, expressed in the same units as your Y values.

### 3.6.12.8 Equation: Plateau followed by one phase association

**Introduction**

In the standard one-phase association equation, the increase starts at time zero. This alternative equation is used when you measure a baseline
for a while, then do some experimental intervention that starts the association at some time $X_0$.

**Entering data**

Create an XY data table. Enter time into $X$, and response (binding, concentration ..) into $Y$. If you have several experimental conditions, place the first into column A, the second into column B, etc.

After entering data, click Analyze, choose nonlinear regression, choose the panel of exponential equations, and choose *Plateau followed by one phase association*.

**Consider constraining $X_0$ to a constant values**

If you know the time at which you initiated the association, you should constrain $X_0$ to that value. To do this, go to the Constrain tab of the nonlinear regression dialog, set the drop down next to $X_0$ to "Constant equal to" and enter the value.

**Model**

$$Y = \begin{cases} X<X_0, & Y_0, \\ Y_0 + (\text{Plateau}-Y_0) \times (1 - \exp(-K \times (X-X_0))) & \end{cases}$$

$X_0$ is the time at which the association begins. Often you will set that to a constant value based on your experimental design, but otherwise Prism can fit it. It is expressed in the same time units as $X$. 
YO is the average Y value up to time X0. It is expressed in the same units as Y.

Plateau is the Y value at infinite times, expressed in the same units as Y.

K is the rate constant, expressed in reciprocal of the X axis time units. If X is in minutes, then K is expressed in inverse minutes.

Tau is the time constant, expressed in the same units as the X axis. It is computed as the reciprocal of K.

Half-life is in the time units of the X axis. It is computed as ln(2)/K.

Span is the difference between Y0 and Plateau, expressed in the same units as your Y values.

3.6.12.9 Equation: Two phase association

Introduction

An exponential decay equation models many chemical and biological processes. It is used whenever the rate at which something happens is proportional to the amount which is left.

A two-phase model is used when the outcome you measure is the result of the sum of a fast and slow exponential decay.

Entering data

Create an XY data table. Enter time into X, and response (binding, concentration ..) into Y. If you have several experimental conditions, place the first into column A, the second into column B, etc.

After entering data, click Analyze, choose nonlinear regression, choose the panel of exponential equations, and choose Two phase association.

Consider constraining Plateau to a constant value of zero

If you have subtracted off any background signal, then you know the curve has to plateau at Y=0. In this case, you should constrain the
parameter Plateau to be a constant value equal to zero. To do this, go to
the Constrain tab of the nonlinear regression dialog, set the drop down
next to Plateau to "Constant equal to" and enter the value 0.0.

Model

\[
\begin{align*}
\text{SpanFast} &= (\text{Plateau} - Y_0) \times \text{PercentFast} \times 0.01 \\
\text{SpanSlow} &= (\text{Plateau} - Y_0) \times (100 - \text{PercentFast}) \times 0.01 \\
Y &= Y_0 + \text{SpanFast} \times (1 - \exp(-K_{\text{Fast}} \times X)) + \text{SpanSlow} \times (1 - \exp(-K_{\text{Slow}} \times X))
\end{align*}
\]

\( Y_0 \) is the \( Y \) value when \( X \) (time) is zero. It is expressed in the same units as \( Y \).

\( \text{Plateau} \) is the \( Y \) value at infinite times, expressed in the same units as \( Y \).

\( K_{\text{Fast}} \) and \( K_{\text{Slow}} \) are the two rate constant, expressed in reciprocal of
the \( X \) axis time units. If \( X \) is in minutes, then \( K \) is expressed in inverse
minutes.

\( \text{Tau}_{\text{Fast}} \) and \( \text{Tau}_{\text{Slow}} \) are the two time constants, expressed in the
same units as the \( X \) axis. They are computed as the reciprocals of the
rate constants.
**Half-time (fast)** and **Half-time (slow)** are in the time units of the X axis. They are computed as \( \ln(2)/K \).

**PercentFast** is the fraction of the span (from Y0 to Plateau) accounted for by the faster of the two components.

### 3.6.12.10 Equation: Exponential growth

#### Introduction

This equation describes the growth with a constant doubling time.

#### Entering data

Create an XY data table. Enter time into X, and response (cell number .) into Y. If you have several experimental conditions, place the first into column A, the second into column B, etc.

Note that Y values must be the actual values. If you transform all the values to logarithms, then it rarely would make sense to fit this equation.

After entering data, click Analyze, choose nonlinear regression, choose the panel of exponential equations, and choose *Exponential growth*.

#### Consider constraining Y0 to a constant value

The parameter Y0 is the Y value at time zero. In many cases, you will know this value precisely. If so, you should constrain that parameter to be a constant value. To do this, go to the Constrain tab of the nonlinear regression dialog, set the drop down next to Y0 to "Constant equal to" and enter its value.
Model

\[ Y = Y_0 \exp(k \cdot X) \]

\( Y_0 \) is the Y value when X (time) is zero. It is expressed in the same units as Y,

**K** is the rate constant, expressed in reciprocal of the X axis time units. If X is in minutes, then K is expressed in inverse minutes.

**Tau** is the time constant, expressed in the same units as the X axis. It is computed as the reciprocal of K.

**Doubling-time** is in the time units of the X axis. It is computed as \( \ln(2)/K \).

### 3.6.13 Lines

#### 3.6.13.1 Key concepts: Fitting lines

**Choosing nonlinear regression, rather than linear regression to fit a line**
Prism offers separate analyses for linear regression and nonlinear regression. But the nonlinear regression analysis can fit a straight-line model. This is useful when you want to take advantage of features in Prism’s nonlinear regression analysis that Prism does not offer in its linear regression analysis, such as the ability to compare two models, apply weighting, or automatically exclude outliers. See a longer discussion of the advantages of using the nonlinear regression analysis to fit a straight line.

Fitting straight lines on graphs with nonlinear axes

The nonlinear regression analysis fits the data, not the graph. Since Prism lets you choose logarithmic or probability axes, some graphs with data points that form a straight line follow nonlinear relationships. Prism’s collection of "Lines" equations includes those that let you fit nonlinear models to graphs that appear linear when the X axis is logarithmic, the Y axis is logarithmic, both axes are logarithmic, or when the Y axis uses a probability scale. In these cases, linear regression will fit a straight line to the data but the graph will appear curved since an axis (or both axes) are not linear. In contrast, nonlinear regression to an appropriate nonlinear model will create a curve that appears straight on these axes.

Segmental linear regression

Segmental regression fits one line to all data points with X less than some value X0, and another line to all points with X greater than X0, ensuring that the two lines intersect at X0.

Segmental linear regression is helpful when X is time, and you did something at time=X0 to change the slope of the line. Perhaps you injected a drug, or rapidly changed the temperature. In these cases, your model really does have two slopes with a sharp transition point.

3.6.13.2 Equation: Fitting a straight line with nonlinear regression

Introduction

Linear regression fits a straight line through your data. Nonlinear regression fits any model, which includes a straight line model. Prism offers separate analyses for linear regression and nonlinear regression, so you can choose either one to fit a line.
Prism's nonlinear regression analysis offers more options than its linear regression analysis, such as the ability to compare two models, apply weighting, automatically exclude outliers and perform normality tests on the residuals. See a longer discussion of the advantages of using the nonlinear regression analysis to fit a straight line.

Step by step

Create an XY data table. There is one X column, and many Y columns. If you have several experimental conditions, place the first into column A, the second into column B, etc.

After entering data, click Analyze, choose nonlinear regression, choose the panel equations for lines, and choose Straight line.

Model

\[ Y = Y_{\text{Intercept}} + \text{Slope} \times X \]

Interpret the parameters

*Y_{\text{Intercept}}* is the Y value where the line intersects the Y axis.

*Slope* is the slope of the line, expressed in Y units divided by X units.

Special forms of the linear regression equation

Horizontal line

If you constrain the slope to be zero, the line will be horizontal. The only parameter is the Y intercept. Prism has this model built in as "horizontal line". The best-fit value of the Y intercept is the mean of all the Y values. The model is:

\[ Y = \text{Mean} + 0 \times X \]

Prism requires that all equations include X. Here X is multiplied by zero, so it is present (as required) but has no effect.
**Line through origin**

If you constrain the Y intercept to be zero, the line has to go through the origin (X=0, Y=0). Prism has this "Line through origin" model built in:

\[ Y = \text{Slope} \times X \]

The only parameter is the slope.

### 3.6.13.3 Equation: Line through origin

**Introduction**

Prism's linear regression analysis fits a straight line through your data, and lets you force the line to go through the origin. This is useful when you are sure that the line must begin at the origin (X=0 and Y=0).

Prism's nonlinear regression offers the equation Line through origin. It offers more options than its linear regression analysis, such as the ability to compare two models, apply weighting, automatically exclude outliers, and perform normality tests on the residuals. See a longer discussion of the advantages of using the nonlinear regression analysis to fit a straight line.

**When to force the line through the origin**

In many scientific situations, it just makes sense that when X=0, Y must also equal 0, so the line should be forced to go through the origin (X=0, Y=0). But even in these situations, it can make sense to fit an ordinary linear regression line that also fits the intercept. The data you are analyzing may be far from the origin, and you may get a better fit through the points (what you care about) when you don't force the line to go through the origin. This can happen when the true model is curved (beginning at the origin), so a line through the data points that is not forced to the origin may fit a whole lot better than line forced to go through the origin.

Prism makes it easy to compare the fit that goes through the origin with one that doesn't.
Step by step

Create an XY data table. There is one X column, and many Y columns. If you have several experimental conditions, place the first into column A, the second into column B, etc.

After entering data, click Analyze, choose nonlinear regression, choose the panel equations for lines, and choose Line Through Origin

Model

\[ Y = \text{Slope} \times X \]

Slope is the slope of the line, expressed in Y units divided by X units. It estimates the ratio of \( Y/X \) in the entire population.

Weighting

In situations where linear regression through the origin is appropriate, it is common for the variation among replicate Y values increases as X (and Y) increase. Prism provides two weighting choices (in the Weights tab) that are used in this situation. Weight by \( 1/X^2 \) when you think the variance in Y is proportional to the square of X, which means the SD among Y values is proportional to X. Weight by \( 1/X \) if you think the variance in Y is proportional to X.

\( R^2 \) when fitting a line through the origin

When you constrain a line to go through a point, there are two possible ways to compute \( R^2 \):

- Compare the fit of the best-fit line with the fit of a horizontal line at the mean Y value. But that null hypothesis (horizontal line through the Ymean) doesn't obey the constraint that it go through the origin.

- Compare the best-fit line with a horizontal line at \( Y=0 \). This obeys the constraint, but often fits the data really badly, pushing up the \( R^2 \) value.

When you use nonlinear regression to fit a line through the origin, Prism uses the first definition above. If you use linear regression, and ask Prism to constrain the line to go through the origin, it simply will not report \( R^2 \) at all.
Reference


3.6.13.4 Equation: Segmental linear regression

**Introduction**

Segmental regression fits one line to all data points with X less than some value X0, and another line to all points with X greater than X0, while ensuring that the two lines intersect at X0.

Segmental linear regression is helpful when X is time, and you did something at time=X0 to change the slope of the line. Perhaps you injected a drug, or rapidly changed the temperature. In these cases, your model really does have two slopes with a sharp transition point.

In other cases, the true model has the slope gradually changing. The data fit a curve, not two straight lines. In this situation, fitting the data with segmental linear regression is not helpful.

**Caution**

Don't use segmental linear regression to analyze a biphasic Scatchard or Lineweaver-Burk plot. A biphasic Scatchard plot follows a curve, not two intersecting lines. There is no abrupt break point. You should fit the original data to a two-site binding curve instead.

**Step by step**

Create an XY data table. Enter time into X, and your measurements into Y. If you have several experimental conditions, place the first into column A, the second into column B, etc.

After entering data, click Analyze, choose nonlinear regression, choose the panel equations for lines, and choose *Segmental linear regression*.

**Model**

\[
Y_1 = \text{intercept1} + \text{slope1} \times X \\
Y_\text{atX0} = \text{slope1} \times X_0 + \text{intercept1} \\
Y_2 = Y_\text{atX0} + \text{slope2} \times (X - X_0) \\
Y = \text{IF}(X < X_0, Y_1, Y_2)
\]
The first line of the equation defines the first line segment from its intercept and slope.

The second line of the equation computes the Y value of the first regression at the right end of that segment, when X=X0.

The third line of the equation computes the second regression segment. Since we want a continuous line, the Y value at the left end of the second segment must equal the Y value at the right end of the first segment (YatX0). The Y value at any other position along the second segment equals YatX0 plus the increase due to the second regression line. That increase equals the slope of the second segment (slope2) times the distance from X0 to X.

The final line defines Y for all values of X. If X is less than X0 then Y is set equal to Y1. Otherwise Y is set equal to Y2.

Interpret the parameters

**Intercept1** is the Y value where the first line segment intersects the Y axis.

**Slope1** is the slope of the first line segment, expressed in Y units divided by X units.

**Intercept2** is the Y value where the second (rightmost) line segment intersects the Y axis.

**Slope2** is the slope of the second line segment, expressed in Y units divided by X units.
**X0** is the X value where the two line segments intersect. Often you will want to constrain this to a constant value equal to the time you applied an experimental intervention.

**Extending to three segments**

Prism does not include an equation for segmental regression with three segments, but you could enter this equation as an user-defined equation:

\[
\begin{align*}
Y_1 &= \text{intercept}_1 + \text{slope}_1 \times X \\
Y_{atX0} &= \text{intercept}_1 + \text{slope}_1 \times X_0 \\
Y_2 &= Y_{atX0} + \text{slope}_2 \times (X - X_0) \\
Y_{atX1} &= Y_{atX0} + \text{slope}_2 \times (X_1 - X_0) \\
Y_3 &= Y_{atX1} + \text{slope}_3 \times (X - X_1) \\
Y &= \text{IF}(X < X_0, Y_1, \text{IF}(X < X_1, Y_2, Y_3))
\end{align*}
\]

3.6.13.5 **Equation: Fitting a straight line on a semi-log or log-log graph**

**Straight lines on graphs with logarithmic axes**

The nonlinear regression analysis fits the data, not the graph. Since Prism lets you choose logarithmic axes, some graphs with data points that form a straight line follow nonlinear relationships. Prism's collection of "Lines" equations includes those that let you fit nonlinear models to graphs that appear linear when the X axis is logarithmic, the Y axis is logarithmic, or both axes are logarithmic. In these cases, linear regression will fit a straight line to the data but the graph will appear curved since an axis (or both axes) are not linear. In contrast, nonlinear regression to an appropriate nonlinear model will create a curve that appears straight on these axes.

**Entering and fitting data**

1. Create an XY table, and enter your X and Y values.
2. Go to the graph, double click on an axis to bring up the Format Axis dialog. Change one or both axes to a logarithmic scale.

3. Click Analyze, choose Nonlinear regression (not Linear regression) and then choose one of the semi-log or log-log equations from the "Lines" section of equations.

**Equations**

**Semilog line -- X axis is logarithmic, Y axis is linear**

\[ Y = \text{Yintercept} + \text{Slope} \times \log(X) \]

**On semilog axis**

Slope is the change in Y when the log(X) changes by 1.0 (so X changes by a factor of 10).

Yintercept is the Y value when log(X) equals 0.0. So it is the Y value when X equals 1.0.
Semilog line -- X axis is linear, Y axis is logarithmic

\[ Y = 10^{(\text{Slope} \times X + \text{Yintercept})} \]

On semilog axis

On linear axes

Slope is the change in \( \log(Y) \) when the \( X \) changes by 1.0.

Yintercept is the \( \log(Y) \) value when \( X \) equals 0.0.

Log-log line -- Both \( X \) and \( Y \) axes are logarithmic

\[ Y = 10^{(\text{slope} \times \log(X) + \text{Yintercept})} \]

On log-log axes

On linear axes
Since both axes are transformed the same way, the graph is linear on both sets of axes. But when you fit the data, the two fits will not be quite identical.

Slope is the change in log(Y) when the log(X) changes by 1.0.

Yintercept is the Y value when log(X) equals 0.0. So it is the Y value when X equals 1.0.

**An alternative way to handle these data**

The nonlinear regression analysis minimizes the sum of the squares of the difference between the actual Y value and the Y value predicted by the curve. This is not the same as minimizing the sum of squares of the distances (as seen on the graph) between points and curve. In the graph below, the two vertical lines look the same distance but one represents a difference of 9 Y units, and the other a difference of 900.
An alternative approach, that might be better in some circumstances, is to use Prism's transform analysis to transform Y (and maybe also X) to logarithms. Then perform linear regression on the logarithms. The regression results will not be the same as using nonlinear regression on log axes.

3.6.13.6 Equation: Fitting a straight line on a graph with a probability axis

Fitting straight lines on graphs with nonlinear axes

The nonlinear regression analysis fits the data, not the graph. If you choose probability Y axis, graphs with data points that form a straight line follow nonlinear relationships. Prism's collection of "Lines" equations includes those that let you fit nonlinear models to graphs that appear linear when the Y axis is a probability axis. In these cases, linear regression will fit a straight line to the data but the graph will appear curved since an axis (or both axes) are not linear. In contrast, nonlinear regression to an appropriate nonlinear model will create a curve that appears straight on these axes.

Entering and fitting data

1. Create an XY table, and enter your X and Y values.
2. Go to the graph, double click on an axis to bring up the Format Axis dialog. Change the Y axis to a probability scale.

3. Click Analyze, choose Nonlinear regression (not Linear regression) and then choose one of the Cumulative Gaussian distribution equations from the "Lines" section of equations.

Equations

**Cumulative Gaussian - Y values are percentages**

Top = 100  
$z = \frac{X - \text{Mean}}{\text{SD}}$  
$Y = \text{Top} \times \text{zdist}(z)$

**On probability axis**

![Probability axis graph]

**On linear axis**

![Linear axis graph]

**Cumulative Gaussian - Y values are fractions**

Top = 1  
$z = \frac{X - \text{Mean}}{\text{SD}}$  
$Y = \text{Top} \times \text{zdist}(z)$
Parameters

**Mean** is the average of the original distribution, from which the frequency distribution was created.

**SD** is the standard deviation of the original distribution.

Both of these parameters are expressed in the same units as the X values plotted on the graph, which is the same as the Y values in the original distribution from which the frequency distribution was generated.

### 3.6.14 Polynomial

#### 3.6.14.1 Key concepts: Polynomial

**Usefulness of polynomial models**

There are two situations where you might want to choose a polynomial model:

- Your scientific model is described by a polynomial equation. This is rare in biology. Few chemical or pharmacological models are described by polynomial equations.

- You don't have a scientific model, but want to fit a curve to interpolate unknown values. With this goal, you often don't care much
about the details of the model. Instead, you care only about finding a model that goes near the data points. Polynomial models often work well.

**Which polynomial model?**

The order of a polynomial model expresses how many terms it has. Prism offers up to a sixth order equation (and it would be easier to enter higher order equations). The higher order equations have more inflection points.

Choosing the best polynomial model is often a matter of trial and error. If the curve doesn't follow the trend of your data, pick a higher order equation. If it wiggles too much, pick a lower order equation.

**How are polynomial models special?**

To a mathematician, polynomial models are very special. Strictly speaking, polynomial models are not 'nonlinear'. Even though a graph of X vs. Y is curved (in all but some special cases), the derivative of Y with respect to the parameters is linear.

Because polynomial models are not nonlinear, it is possible (but not with Prism) to fit polynomial models without fussing with initial values. And the fit can be in one step, rather than the iterative approach used for nonlinear models.

Since Prism treats polynomial models the same way it treats nonlinear models, it does require initial values (it chooses 1.0 for each parameter automatically). It doesn't matter what values are used -- polynomial regression cannot encounter false minima.

**Why you should choose a centered polynomial equation**

There are two problems with polynomial fits, often solved by centering:

- When the X values are large, and start well above zero (for example, when X is a calendar year), taking the very large X values to large powers can lead to math errors.
- Even when the X values are not large, the parameters of the model are intertwined, so have high covariance and dependency. This results in large standard errors, wide confidence intervals, and huge confidence or prediction bands.

Both these problems are solved by using a centered polynomial model. The idea of centering is simple. Subtract the mean of all X values from each X value, and use those differences instead of X in the model.
Fitting the centered model leads to exactly the same curve (unless the regular approach led to math errors). Accordingly, the sum-of-squares are $R^2$ are the same, as are results of model comparisons. However, the parameters have different meanings, so have different best-fit values (except the first parameter which is the same), different standard errors and confidence intervals, smaller covariances and dependencies, and tighter confidence/prediction bands.

Read more about centered polynomial models.

### 3.6.14.2 Centered polynomial equations

Included in Prism, are both a set of ordinary polynomial equations and also a set of centered polynomial equations. For example, when you look in the list of polynomials you'll see both 'Second order polynomial' and 'Centered second order polynomial'. We recommend always choosing one of the centered equations instead of an ordinary polynomial equation. This page explains why.

**What's wrong with ordinary polynomial models?**

The standard polynomial models look like this:

$$Y = B_0 + B_1 \times X + B_2 \times X^2$$

More terms are included with the higher order equations.

There are two problems with polynomial fits:

- When the X values are large, and start well above zero (for example, when X is a calendar year), taking the very large X values to large powers can lead to math overflow. Even if the program doesn't report any math error, the results can be inaccurate. Some coefficients will be positive and some negative, so the value of Y depends on subtracting huge numbers from other huge numbers, leading to imprecise results.

- Even when the X values are not large, the parameters of the model are intertwined, so have high covariance and dependency. This results in large standard errors, wide confidence intervals, and very wide confidence or prediction bands. In many cases, this problem is severe enough that Prism reports that the results are 'ambiguous' and so doesn't report confidence intervals for all the parameters and can't graph confidence bands.
What are centered polynomial models?

Both problems go away when the X values are centered. The idea of centering is to subtract the mean X from all X values before fitting the model. This can be done as part of nonlinear regression, using this model:

\[ XC = X - X_{\text{mean}} \]
\[ Y = B_0 + B_1 \times XC + B_2 \times XC^2 \]

Here XC is the centered X value, equal to the X value minus Xmean, which is the mean of all X values. In other words, XC is the distance of any X value from the mean of all X values. Xmean is constant, and not a parameter that Prism tries to fit. Of course, you can include more terms in the definition of Y to create higher order polynomial equations.

The advantages of centered models

Fitting the centered model leads to exactly the same curve (unless the regular approach led to math errors). Accordingly, the sum-of-squares is the same, as are results of model comparisons.

However, the centered equation has reparameterized the model. The parameters have different meanings, so have different best-fit values (except the first parameter which is the same), different standard errors and confidence intervals, smaller covariances and dependencies, and tighter confidence/prediction bands.

How centered models are implemented in Prism

You can fit data to a built-in centered polynomial equation without knowing how Prism implements the model. If you are curious, read on.

The built-in set of centered polynomial equations, written as shown above, constrain the parameter XMean to equal the mean of X value by constraining it to equal a "Data set constant (= Mean X)".

If you open a file using centered polynomial regression in an version of Prism prior to 5.02 or 5.0b, that constraint will be lost, and centered polynomial regression won't work.
3.6.14.3 Equations: Polynomial models

**Step by step**

Create an XY data table. There is one X column, and many Y columns. If you have several experimental conditions, place the first into column A, the second into column B, etc.

After entering data, click Analyze, choose nonlinear regression, choose the panel equations for polynomial equations, and choose one.

The "order" of a polynomial equation tells you how many terms are in the equation. Prism offers first to sixth order polynomial equations (and you could enter higher order equations as user-defined equations if you need them). Higher order models wiggle more than do lower order models. Since the equation rarely corresponds to a scientific model, use trial and error. If it isn't close enough to the data, pick a higher order equation. If it wiggles too much, pick a lower order equation.

**Polynomial models**

<table>
<thead>
<tr>
<th>Order</th>
<th>Equation</th>
</tr>
</thead>
<tbody>
<tr>
<td>First</td>
<td>( Y = B_0 + B_1 \times X ) (straight line)</td>
</tr>
<tr>
<td>Second</td>
<td>( Y = B_0 + B_1 \times X + B_2 \times X^2 ) (quadratic equation)</td>
</tr>
<tr>
<td>Third</td>
<td>( Y = B_0 + B_1 \times X + B_2 \times X^2 + B_3 \times X^3 )</td>
</tr>
<tr>
<td>Fourth</td>
<td>( Y = B_0 + B_1 \times X + B_2 \times X^2 + B_3 \times X^3 + B_4 \times X^4 )</td>
</tr>
<tr>
<td>Fifth</td>
<td>( Y = B_0 + B_1 \times X + B_2 \times X^2 + B_3 \times X^3 + B_4 \times X^4 + B_5 \times X^5 )</td>
</tr>
<tr>
<td>Sixth</td>
<td>( Y = B_0 + B_1 \times X + B_2 \times X^2 + B_3 \times X^3 + B_4 \times X^4 + B_5 \times X^5 + B_6 \times X^6 )</td>
</tr>
</tbody>
</table>

There is no general way to interpret the coefficients \( B_0, B_1, \) etc. In most cases, the goal of fitting a polynomial model is to make a curve that looks good, and the parameters really don't matter.
**Centered polynomial models**

The **Centered polynomial models** are identical to the ones listed above, with one exception. Wherever X appears above, replace it with \((X - X_{\text{mean}})\), where \(X_{\text{mean}}\) is the mean of all X values (for rows that have Y values). Using a centered model can avoid computer math problems (overflows), and we recommend that you use them routinely.

### 3.6.15 Gaussian

#### 3.6.15.1 Key concepts: Gaussian

**What is a frequency distribution?**

All the equations in this section fit frequency distributions. The X values represent Y values in the original data set, and the Y values are the frequency or cumulative frequency.

Prism can create a frequency distribution from column data, using an analysis created for that purpose (called, of course, Frequency Distribution).

**What is a cumulative frequency distribution?**

A frequency distribution plots the number of observations as a function of value. A cumulative frequency distribution plots the *cumulative* number of observations as a function of value. Each Y value is the number of observations in the original data set that have a value less than or equal to the X value.

The Y values can be expressed as the counted number of observations, as fractions or as percentages.
3.6.15.2 Equation: Gaussian distribution

**Introduction**

Data follow a Gaussian distribution when scatter is caused by the sum of many independent and equally weighted factors.

A frequency distribution (histogram) created from Gaussian data will look like a bell-shaped Gaussian distribution.

**Step-by-step**

The data you fit must be in the form of a frequency distribution on an XY table. The X values are the bin center and the Y values are the number of observations.

If you start with a column of data, and use Prism to create the frequency distribution, make sure that you set the graph type to "XY graph", with either points or histogram spikes. This ensures that Prism creates an XY results table with the bin centers entered as X values. If you pick a bar graph instead, Prism creates a column results table, creating row labels from the bin centers. This kind of table cannot be fit by nonlinear regression, as it has no X values.

Starting from the frequency distribution table, click Analyze, choose Nonlinear regression from the list of XY analyses, and then choose the "Gaussian" equation from the "Gaussian" family of equations.

**Model (Gaussian distribution)**

\[ Y = \text{Amplitude} \times \exp\left(-0.5\times\left(\frac{X-\text{Mean}}{\text{SD}}\right)^2\right) \]
**Amplitude** is the height of the center of the distribution in Y units.

**Mean** is the X value at the center of the distribution.

**SD** is a measure of the width of the distribution, in the same units as X.

The mean and SD will not be identical to the mean and SD computed directly from the raw data. There are two reasons for the discrepancy. The first is that creating the frequency distribution requires a fairly arbitrary decision about bin width, and that will influence the best-fit values of Mean and SD. The second reason is that the nonlinear regression assumes that the residuals (the distances of the points from the curve) follow a Gaussian distribution. This assumption won't be exactly true in a frequency distribution.

**Model (sum of two Gaussian distributions)**

If your data are a mixture of values sampled from two Gaussian distributions, fit the frequency distribution to a model of the sum of two Gaussian distributions.

\[
\text{One} = \text{Amplitude1} \times \exp\left(-0.5 \times \left(\frac{\text{X-Mean1}}{\text{SD1}}\right)^2\right) \\
\text{Two} = \text{Amplitude2} \times \exp\left(-0.5 \times \left(\frac{\text{X-Mean2}}{\text{SD2}}\right)^2\right) \\
Y = \text{One} + \text{Two}
\]

**Amplitude1** and **Amplitude2** are the heights of the center of the distribution in Y units.

**Mean1** and **Mean2** are the X values at the center of the two distributions.
**SD1** and **SD2** are measures of the widths of the distributions, in the same units as X.

Prism is not very smart about assigning initial values to the parameters. If you have trouble getting this model to fit, try fussing with the initial parameter values.

**Area under a Gaussian distribution**

The area under a Gaussian distribution equals Amplitude*SD/0.3989. That constant equals the reciprocal of the square root of two pi.

### 3.6.15.3 Equation: Log Gaussian distribution

Note: Versions of Prism up to 7.00 and 7.0a used a different and nonstandard form of this equation which we called log(Gaussian).

**Introduction**

Data follow a Gaussian distribution when scatter is caused by the **sum** of many independent and equally weighted factors.

When scatter is caused by the **product** of many independent and equally weighted factors, data follow a lognormal distribution. When plotted on a linear X axis, this is skewed to the right (see below). When plotted on a logarithmic X axis, it looks like a bell-shaped Gaussian distribution.

**Step-by-step**

The data must be in the form of a frequency distribution on an XY table. The X values are the bin center and the Y values are the number of observations.

If you start with a column of data, and use Prism to create the frequency distribution, make sure that you set the graph type to "XY graph", with either points or histogram spikes. This ensures that Prism creates an XY results table with the bin centers entered as X values. If you pick a bar graph instead, Prism creates a column results table, creating row labels from the bin centers. This kind of table cannot be fit by nonlinear regression, as it has no X values.
Starting from the frequency distribution table, click Analyze, choose Nonlinear regression from the list of XY analyses, and then choose the "lognormal" equation from the "Gaussian" family of equations.

**Model**

\[ Y = \frac{A}{X} \exp\left(-0.5 \left( \ln\left(\frac{X}{\text{GeoMean}}\right) / \ln(\text{GeoSD}) \right)^2 \right) \]

**GeoMean** is the geometric mean in the units of the data.

**GeoSD** is the geometric standard deviation factor. It is a unitless ratio.

**A** is related to the amplitude and area of the distribution.

\[
\text{Amplitude} = \frac{A}{\left(\text{GeoMean} / \exp(0.5\ln(\text{GeoSD})^2)\right)}
\]

\[
\text{Area} = A \times \sqrt{2 \times \pi} \times \ln(\text{GeoSD})
\]

A more standard form of the model (from Wikipedia or MathWorld) is:

\[ Y = \frac{1}{X \times S \times \sqrt{2 \times \pi}} \exp\left(-0.5 \left( \ln(X) - M \right)^2 / (S^2) \right) \]

This standard form of the equation doesn't have the A parameter because the area under the curve, using the equation above, is always 1.0. S in this model equals \ln(\text{GeoSD}) and M equals \ln(\text{GeoMean}).
3.6.15.4 Equation: Cumulative Gaussian distribution

**Introduction**

A frequency distribution plots the number of observations as a function of value. A cumulative frequency distribution plots the *cumulative* number of observations as a function of value. Each Y value is the number of observations in the original data set that have a value less than or equal to the X value.

The advantage of creating a cumulative distribution is that you don't have to make any choice regarding bin width.

If your data follow a Gaussian distribution, the cumulative distribution has a sigmoidal shape.

![Cumulative Gaussian distribution graph]

**Step-by-step**

1. Create an XY table, and enter your X and Y values. The X values correspond to the value in the original data set, and the Y values are the number (or fraction or percent) of values in the original data set that are less than or equal to the Y value.

   Alternatively, enter a stack of values onto a Column data table, and run the frequency distribution analysis choosing to create a cumulative frequency distribution with no bins.

2. From the cumulative frequency distribution, click Analyze, choose Nonlinear regression and then choose one of the Cumulative...
Gaussian distribution equations from the "Gaussian" group of equations.

3. If your data are entered as counts (rather than percentages or fractions) constrain N to a constant value equal to the number of observations.

Models

The details of the model depend on whether the Y values are percentages, fractions or counts.

Here is the model if the data are percentages, so the last Y value equals 100.

\[ \text{Top} = 100 \\
\text{z} = \frac{(X - \text{Mean})}{\text{SD}} \\
\text{Y} = \text{Top} \times \text{zdist}(z) \]

Here is the model if the data are fractions, so the first line of the model defines Top to equal 1.00.

\[ \text{Top} = 1.0 \\
\text{z} = \frac{(X - \text{Mean})}{\text{SD}} \\
\text{Y} = \text{Top} \times \text{zdist}(z) \]

And finally, here is the model if the data are numbers of observations, so the largest value equals the number of observations (N). In this case, you should constrain N to be a constant value equal to the number of observations.

\[ \text{z} = \frac{(X - \text{Mean})}{\text{SD}} \\
\text{Y} = N \times \text{zdist}(z) \]

**Mean** is the average of the original distribution, from which the frequency distribution was created.

**SD** is the standard deviation of the original distribution.

Both of these parameters are expressed in the same units as the X values plotted on the graph, which is the same as the Y values in the original distribution from which the frequency distribution was generated.
Plotting on a log Y axis

If you choose a Y axis with a probability scale, then the cumulative Gaussian distribution appears as a straight line. For this reason, the cumulative Gaussian models are part of both "Lines" families of equations as well as the "Gaussian" family. The two listings are identical.

3.6.15.5 Equation: Lorentzian

Introduction

A Lorentzian distribution is bell shaped, but has much wider tails than does a Gaussian distribution.

Step-by-step

The data must be in the form of a frequency distribution on an XY table. The X values are the bin center and the Y values are the number of observations.

If you start with a column of data, and use Prism to create the frequency distribution, make sure that you set the graph type to "XY graph", with either points or histogram spikes. This ensures that Prism creates an XY results table with the bin centers are entered as X values. If you pick a bar graph instead, Prism creates a column results table, creating row labels from the bin centers. This kind of table cannot be fit by nonlinear regression, as it has no X values.

Starting from the frequency distribution table, click Analyze, choose Nonlinear regression from the list of XY analyses, and then choose the "Lorentzian" equation from the "Gaussian" family of equations.
Model (Lorentzian distribution)

\[ Y = \frac{\text{Amplitude}}{1 + ((X - \text{Center}) / \text{Width})^2} \]

**Amplitude** is the height of the center of the distribution in Y units.

**Center** is the X value at the center of the distribution.

**Width** is a measure of the width of the distribution, in the same units as X. This is not identical to a standard deviation, but has the same general meaning.

Model (sum of two Lorentzian distributions)

\[ \begin{align*} 
\text{One} &= \frac{\text{Amplitude}_1}{1 + ((X - \text{Center}_1) / \text{Width}_1)^2} \\
\text{Two} &= \frac{\text{Amplitude}_2}{1 + ((X - \text{Center}_2) / \text{Width}_2)^2} \\
Y &= \text{One} + \text{Two} 
\end{align*} \]

**Amplitude_1** and **Amplitude_2** are the heights of the center of the distribution in Y units.

**Center_1** and **Center_2** are the X values at the center of the two distributions.

**Width_1** and **Width_2** are measures of the widths of the distributions, in the same units as X.

Prism is not very smart about assigning initial values to the parameters. If you have trouble getting this model to fit, try fussing with the initial parameter values.

3.6.16 Sine waves

3.6.16.1 Standard sine wave

**Introduction**

Sine waves describe many oscillating phenomena.
Step by step

Create an XY data table. There is one X column, and many Y columns. If you have several experimental conditions, place the first into column A, the second into column B, etc.

After entering data, click Analyze, choose nonlinear regression, choose the panel of equations for sine waves, and choose Standard sine wave.

If you know the Y value must equal zero at time zero, then constrain PhaseShift to a constant value of zero.

You may need to fuss with the initial values for PhaseShift and Wavelength, as our built-in rules for computing the initial value rarely work.

- For Wavelength. Count the number of peaks between two values X = a and X = b and set the initial value for Wavelength to \( \frac{1}{\left( \text{NumberOfPeaks} \div (b - a) \right)} \).

- For PhaseShift. If Y=0 at X=0, then PhaseShift is zero. If Y is at its maximum when X=0, then PhaseShift=\( \pi \) (3.14159). Interpolate in between if necessary.

Model

\[
Y = \text{Amplitude}\sin(\frac{2\pi X}{\text{Wavelength}} + \text{PhaseShift})
\]

Interpret the parameters

**Amplitude** is the height of top of the waves, in Y units. If Prism reports a negative Amplitude but you expect a positive Amplitude, simply change the initial value of the PhaseShift by \( \pi \) in either direction.
**Wavelength** is the time it takes for a complete cycle, in units of X.

**Frequency** is the number of cycles per time unit. It is calculated as the reciprocal of wavelength, and is expressed in the inverse of the time units of X.

**PhaseShift** in radians. A PhaseShift of 0 sets Y equal to 0 at X=0. A PhaseShift of π sets Y equal to its maximum when X=0.

### 3.6.16.2 Damped sine wave

**Introduction**

Sine waves describe many oscillating phenomena. Often the peak of each wave decreases or dampens as time goes on.

**Step by step**

Create an XY data table. There is one X column, and many Y columns. If you have several experimental conditions, place the first into column A, the second into column B, etc.

After entering data, click Analyze, choose nonlinear regression, choose the panel of equations for sine waves, and choose *Damped sine wave*.

If you know the Y value is zero at time zero, then constrain PhaseShift to a constant value of zero.

You may need to fuss with the initial values for PhaseShift and Wavelength, as our built-in rules for computing the initial values don't always work very well.

**Model**

\[ Y = \text{Amplitude}\times \exp(-K\times X)\times \sin\left((2\times \pi \times X)/\text{Wavelength}\right)\times \text{PhaseShift} \]
Interpret the parameters

**Amplitude** is the height of top of the waves, in Y units.

**Wavelength** is the time it takes for a complete cycle, in units of X.

**Frequency** is the number of cycles per time unit. It is calculated as the reciprocal of wavelength, and is expressed in the inverse of the time units of X.

**PhaseShift** in radians. A phaseshift of 0 sets Y equal to 0 at X=0.

K is the decay constant, in the reciprocal of the time units of the X axis.

**HalfLlife** is the time it takes for the maximum amplitude to decrease by a factor of 2. It is computed as 0.693/K.

### 3.6.16.3 Sinc wave

**Introduction**

The sinc() function appears frequently in signal and image processing because it is the Fourier transform of a rectangular pulse. It is also called the "sampling" or "sine cardinal" function.

**Step by step**

Create an XY data table. There is one X column, and many Y columns. If you have several experimental conditions, place the first into column A, the second into column B, etc.
After entering data, click Analyze, choose nonlinear regression, choose the panel of equations for sine waves, and choose Sinc() function.

You may need to fuss with the initial values for Phaseshift and Wavelength, as our built-in rules for computing the initial value doesn't always work very well.

**Model**

\[ Y = \begin{cases} 
\text{Amplitude} & \text{if } X = 0 \\
\frac{\text{Amplitude} \cdot \sin\left(\frac{2\pi X}{\text{Wavelength}}\right)}{\frac{2\pi X}{\text{Wavelength}}} & \text{else} 
\end{cases} \]

**Interpret the parameters**

- **Amplitude** is the height of top of the waves, in Y units.
- **Wavelength** is the time it takes for a complete cycle, in units of X
- **Frequency** is the number of cycles per time unit. It is calculated as the reciprocal of wavelength, and is expressed in the inverse of the time units of X.

### 3.6.16.4 Sine wave with nonzero baseline

**Introduction**

Sine waves describe many oscillating phenomena. Usually the sine wave oscillates around \( Y=0 \) (what we call the standard sine wave). This equation has the wave oscillating around a baseline value that does not have to be zero.
Step by step

Create an XY data table. There is one X column, and many Y columns. If you have several experimental conditions, place the first into column A, the second into column B, etc.

After entering data, click Analyze, choose nonlinear regression, choose the panel of equations for sine waves, and choose *Sine wave with nonzero baseline*.

If you know the Y value is at baseline at time zero, then constrain PhaseShift to a constant value of zero.

You may need to fuss with the initial values for PhaseShift and Wavelength, as our built-in rules for computing the initial value rarely work.

- For Wavelength. Count the number of peaks between two values \( X = a \) and \( X = b \) and set the initial value for Wavelength to \( 1 / ( \text{NumberofPeaks} / (b - a)) \).

- For PhaseShift. If \( Y = \text{Baseline} \) at \( X = 0 \), then PhaseShift is zero. If \( Y \) is at its maximum when \( X = 0 \), then PhaseShift = \( \pi \) (3.14159). Interpolate in between if necessary.

Model

\[
Y = \text{Amplitude} \times \sin\left(\frac{2\pi X}{\text{Wavelength}} + \text{PhaseShift}\right) + \text{Baseline}
\]
**Interpret the parameters**

**Amplitude** is the height of top of the waves from the baseline, in Y units. If Prism reports a negative Amplitude but you expect a positive Amplitude, simply change the initial value of the PhaseShift by Pi in either direction.

**Wavelength** is the time it takes for a complete cycle, in units of X.

**Frequency** is the number of cycles per time unit. It is calculated as the reciprocal of wavelength, and is expressed in the inverse of the time units of X.

**PhaseShift** in radians. A phase shift of 0 sets Y equal to Baseline at X=0. A PhaseShift of n sets Y equal to its maximum when X=0.

**Baseline** is the Y value around which the curve oscillates, so is in the same units as Y.

### 3.6.17 Classic equations from prior versions of Prism

#### 3.6.17.1 Equation: One site binding (hyperbola)

\[
Y = \frac{B_{\text{max}} \cdot X}{K_d + X}
\]

This curve is known as a *rectangular hyperbola*, *binding isotherm*, or *saturation binding curve*. Y is zero initially, and increases to a maximum plateau value $B_{\text{max}}$.

This equation describes the equilibrium binding of a ligand to a receptor as a function of increasing ligand concentration.
- X is the concentration of the ligand.

- Y is the specific binding.

- Bmax is the maximum number of binding sites, expressed in the same units as the Y-axis (usually radioactive counts per minute, sites per cell, or fmol of receptor per mg of tissue).

- Kd is the equilibrium dissociation constant, expressed in the same units as the X-axis (concentration). When the drug concentration equals Kd, half the binding sites are occupied at equilibrium.

This equation also describes the activity of an enzyme as a function of substrate concentration. In this case, the variable labeled Bmax is really Vmax, the maximum enzyme activity, and the variable labeled Kd is really Km, the Michaelis-Menten constant.

3.6.17.2 Equation: Two site binding

\[ Y = \frac{B_{\text{max}1} \cdot X}{K_{d1} + X} + \frac{B_{\text{max}2} \cdot X}{K_{d2} + X} \]

This equation is an extension of the one site binding curve. It shows the binding of a ligand to two receptors with different affinities (different Kd values). It also describes the enzyme activity as a function of substrate concentration when two isozymes are present. The curve in the example has Kd values that differ by a factor of ten, with equal Bmax values. Even with such a large difference between Kd values, the curve is not obviously biphasic.
3.6.17.3 Equation: Sigmoidal dose-response

\[ Y = \text{Bottom} + \frac{(\text{Top} - \text{Bottom})}{1 + 10^{\text{LogEC50} - X}} \]

This is a general equation for a dose-response curve. It shows response as a function of the logarithm of concentration. \( X \) is the logarithm of agonist concentration and \( Y \) is the response. This equation is also called a three-parameter logistic equation.

The variable \( \text{Bottom} \) is the \( Y \) value at the bottom plateau; \( \text{Top} \) is the \( Y \) value at the top plateau, and \( \text{LogEC50} \) is the \( X \) value when the response is halfway between \( \text{Bottom} \) and \( \text{Top} \). \( \text{LogEC50} \) is the logarithm of the EC50 (effective concentration, 50%). With different kinds of variables, this variable is sometimes called \( \text{ED50} \) (effective dose, 50%), or \( \text{IC50} \) (inhibitory concentration, 50%, used when the curve goes downhill).

This equation assumes a standard slope, where the response goes from 10% to 90% of maximal as \( X \) increases over about two log units. The next equation allows for a variable slope.

3.6.17.4 Equation: Sigmoidal dose-response (variable slope)

\[ Y = \text{Bottom} + \frac{(\text{Top} - \text{Bottom})}{1 + 10^{\text{LogEC50} - X \cdot \text{HillSlope}}} \]
This equation extends the previous equation, but allows for a variable slope. This equation is also called a four-parameter logistic equation.

- Bottom is the Y value at the bottom plateau.

- Top is the Y value at the top plateau.

- LogEC50 is the X value when the response is halfway between Bottom and Top. With different kinds of variables, this variable is sometimes called ED50 (effective dose, 50%), or IC50 (inhibitory concentration, 50%, used when the curve goes downhill).

- HillSlope describes the steepness of the curve. This variable is called the Hill slope, the slope factor, or the Hill coefficient. If it is positive, the curve increases as X increases. If it is negative, the curve decreases as X increases. A standard sigmoid dose-response curve (previous equation) has a Hill Slope of 1.0. When HillSlope is less than 1.0, the curve is more shallow. When HillSlope is greater than 1.0, the curve is steeper. The Hill slope has no units.

3.6.17.5 Equation: One site competition

\[ Y = \text{Bottom} + \frac{(\text{Top} - \text{Bottom})}{1 + 10^{X - \text{LogEC50}}} \]

This equation describes the competition of a ligand for receptor binding. It is identical to the sigmoid dose-response curve with \( \text{HILLSLOPE} = -1.0 \).

The variable LogEC50 is the concentration of the competitor required to compete for half the specific binding. We use the term EC50 to be
consistent with the equations for the other sigmoid curves. The term IC50 is used more frequently ("E" stands for effective; "I" stands for inhibitory).

Usually the Y values are total binding. If you enter specific binding instead, fix BOTTOM to have a constant value of zero. If you enter percent specific binding, also set TOP to be a constant equal to 100.

3.6.17.6 Equation: Two site competition

\[
Y = \text{Bottom} + (\text{Top} - \text{Bottom}) \left( \frac{\text{Fraction}_1}{1 + 10^{X - \text{LogEC50}_1}} - \frac{1 - \text{Fraction}_1}{1 + 10^{X - \text{LogEC50}_2}} \right)
\]

This equation describes the competition of a ligand for two types of receptors. The radioligand has identical affinities for both receptors, but the competitor has a different affinity for each.

Y is binding (total or specific) and X is the logarithm of the concentration of the unlabeled ligand. FRACTION_1 is the fraction of the receptors that have an affinity described by LogEC50_1. The remainder of the receptors have an affinity described by LogEC50_2. If LogEC50_1 is smaller than LogEC50_2, then Fraction_1 is the fraction of high affinity sites. If LogEC50_1 is larger than LogEC50_2, then Fraction_1 is the fraction of low affinity sites.
3.6.17.7 Equation: Boltzmann sigmoid

\[
Y = \text{Bottom} + \frac{(\text{Top} - \text{Bottom})}{1 + \exp\left(\frac{V50 - X}{\text{Slope}}\right)}
\]

This equation describes voltage dependent activation of ion channels. It describes conductance (Y) as a function of the membrane potential (X). Conductance varies from BOTTOM to TOP. V50 is the potential at which conductance is halfway between BOTTOM and TOP. SLOPE describes the steepness of the curve, with a larger value denoting a shallow curve. Slope is expressed in units of potential, usually mV, and is positive for channels that activate upon depolarization.

Under appropriate experimental conditions, you can use SLOPE to calculate the valence (charge) of the ion moving across the channel. SLOPE equals RT/zF where R is the universal gas constant, T is temperature in °K, F is the Faraday constant, and z is the valence. Since RT/F \(\approx\) -26 mV at 25°C, \(z = \frac{-26}{\text{SLOPE}}\).

BOTTOM is commonly made a constant equal to 0.0. If you also make TOP a constant equal to 1.0, then Y can be viewed as the fraction of channels that are activated.

3.6.17.8 Equation: One phase exponential decay

\[
Y = \text{Span} \cdot e^{-kX} + \text{Plateau}
\]
This equation describes the kinetics such as the decay of radioactive isotopes, the elimination of drugs, and the dissociation of a ligand from a receptor.

X is time.

Y may be concentration, binding, or response. Y starts out equal to SPAN+PLATEAU and decreases to PLATEAU with a rate constant K.

The half-life of the decay is 0.6932/K.

SPAN and PLATEAU are expressed in the same units as the Y axis. K is expressed in the inverse of the units used by the X axis. In many circumstances, the plateau equals zero. When fitting data to this equation, consider fixing the plateau to a constant value of zero.

3.6.17.9 Equation: Two phase exponential decay

$$Y = Span1 \cdot e^{-K1 \cdot X} + Span2 \cdot e^{-K2 \cdot X} + Plateau$$

This equation describes a two phase exponential decay. Y starts out equal to Span1+Span2+PLATEAU and decays to PLATEAU with fast and slow components. The two half-lives are 0.6932/K1 and 0.6932/K2. In the figure, the two rate constants differ tenfold, but the spans were
equal. The curve is not obviously biphasic, and it takes a very practiced eye to see that the curve does not follow a single phase model.

3.6.17.10 Equation: One phase exponential association

\[ Y = Y_{\text{max}} \left( 1 - e^{-K_{1}X} \right) \]

This equation describes the pseudo-first order association kinetics of the interaction between a ligand and its receptor, or a substrate and an enzyme. Y is either binding or enzyme activity. X is time.

Y starts out equal to zero and increases to a maximum plateau (at equilibrium) equal to YMAX. When X equals 0.6932/K, Y equals 0.5*YMAX.

3.6.17.11 Equation: Two phase exponential association

\[ Y = Y_{\text{max}1} \cdot (1 - e^{-K_{1}X}) + Y_{\text{max}2} \cdot (1 - e^{-K_{2}X}) \]

This is an extension of the exponential association to two phases, corresponding to a radioligand binding to two independent sites.

3.6.17.12 Equation: Exponential growth

\[ Y = \text{Start} \cdot e^{K_{1}X} \]
This describes an exponential growth curve. Y is population size (perhaps cell number) and X is time. At X=0, Y equals START. Y increases geometrically with a doubling time equal to 0.6932/K.

Note: It is difficult to fit data to this equation with nonlinear regression, because a tiny change in the initial values will drastically alter the sum-of-squares. You may need to override the initial values provided by Prism.

3.6.17.13 Equation: Power series

This versatile equation has many uses.

\[ Y = A \cdot X^B + C \cdot X^D \]

Fitting data to a power series model can be difficult. The initial values generated automatically by Prism are not very helpful (all four parameters are set to 1.0). you will probably need to enter better initial values in order to fit this equation to data. The initial values of B and D are important, because small changes in those values can make a huge change in Y.

The equation is not defined, and leads to a floating point error, if X equals zero and B or D are negative numbers or if X is negative and B or D are between 0.0 and 1.0.

3.6.17.14 Equation: Sine wave

\[ Y = \text{Baseline} + \text{Amplitude} \cdot \sin(\text{Frequency} \cdot X + \text{Offset}) \]
X is in radians. In most cases, you will want to fix BASELINE to a constant value of zero. AMPLITUDE is the maximum height of the curve away from the baseline. FREQUENCY is the number of complete oscillations per 1 X unit.

3.6.17.15 Equation: Gaussian distribution

Cumulative probability distribution of a Gaussian bell-shaped distribution with specified mean and SD. The area under the entire curve is AREA. A standard probability distribution is scaled so that AREA equals 1.0. The units of the Y-axis are arbitrary, determined by your choice of AREA.

3.7 Entering a user-defined model into Prism

Prism provides lots of built-in models, but you should not hesitate to adapt these or enter additional models. This section shows you how to do so.

- Overview: User-defined equations
- How to: Enter a new equation
- How to: Clone an equation
- How to: Manage your list of equations
3.7.1 Overview: User-defined equations

Few people will need to enter user-defined equations into Prism. If you need to do this, keep these points in mind:

- Rather than starting with a blank screen, it is usually much easier to clone an existing equation, and then modify it.

- The syntax Prism uses is pretty standard. But use plenty of parentheses, so there is no doubt about what the equation means. Many people have made the mistake of entering "A/B+C" expecting Prism to divide A by the sum of B plus C. Instead, Prism first divides A by B, and then adds that quotient to C. Writing the equation with parentheses as "A/(B+C)" avoids any ambiguity.

- Prism lets you use many standard mathematical functions when defining your model. Most of these are standard. But watch out for a few that are implemented differently in different languages. The log() function in Prism is common (base 10) log. If you want to use
natural logs, use the ln() function. The sqr() function in Prism squares the value, while the sqrt() function computes the square root.

- Prism lets you write an equation in such a way that different models are fit to different data sets.

- When you write a model using several lines, Prism evaluates from the top down. This follows the conventions of all computer languages, but is opposite to how mathematical textbooks present complicated equations.

- After writing the equation, don’t forget to define rules for initial values. If you don’t do this, you’ll need to enter initial estimated values for each parameter every time you fit the model to data.

- When you enter a new equation, Prism also lets you define default constraints and transforms to report. These features are optional, so you can skip them when first learning about user-defined equations.

3.7.2 How to: Enter a new equation

Entering an equation

Prism comes with many built-in equations, but you will often want to enter a different equation. To do so, click the New button on the top of the Fit tab of the Nonlinear regression dialog.

A drop down menu lets you choose to enter a new equation, clone an existing one, or import an equation from a saved .PZF file.
Name, definition, tip and description

Equation type

Choose whether you are entering an ordinary explicit equation (that defines Y as a function of X and parameters), an implicit equation (Y appears on both sides of the equals sign) or a differential equation (which defines the derivative of Y with respect to X).

Name

Enter a name you will recognize, so you can choose the equation again. The name also appears on the results table, so make it both descriptive and concise.

Definition

Write the math that defines Y as a function of X and one or more parameters. Read the details of equation syntax, a list of available
math functions you can use, and how to fit different models to different ranges of X values and to different data sets.

Tip

The tip appears below the equation list in the Fit tab of the Nonlinear regression dialog. Use it to distinguish among similar equations. Tips are optional, and must be short (a few sentences).

Description

The description can be up to several paragraphs. Anyone choosing the equation can view the description (and the math) by clicking the Edit or View button. This is a good place to document the source of the equation, its assumptions, units of the parameters, etc.

Rules for initial values

Nonlinear regression must begin with initial estimated values for each parameter. You can enter values directly, or enter a rule whereby Prism can compute the initial value from the range of the X and Y values. Learn more.

The values or rules you enter will become the default set of initial values that will be used every time the equation is selected. But each time the equation is selected, you (or whoever is selecting the equation) can change the initial values for that one fit.

Default constraints

Use the constraints tab to set default constraints. You can constrain a parameter to a constant value, constrain to a range of values, share among data sets (global fit), or define a parameter to be a column constant. These constraints will become the default every time the equation is selected. But each time the equation is selected, you (or whoever is selecting the equation) can change the constraints for that one fit.

If a parameter has to be set constant, but the actual value is different for each experiment, set the constraint "Constant equal to" but leave the value blank. If someone chooses the equation but forgets to constrain that parameter to a constant value, Prism will prompt for one.
**Transforms to report**

Define transforms of the best-fit values on the Transforms to report tab. Unlike initial values and constraints, you cannot override these transforms each time you choose the equation. Transforms are defined with the equation definition and cannot be tweaked each time the equation is selected.

### 3.7.3 How to: Clone an equation

If none of the built-in equations suit your needs, you may not need to enter a new equation. Instead, consider cloning an existing equation. You may also clone an equation if your only goal is to rename some of the parameters.

**To clone an equation:**

1. Click the New button on the top of the Fit tab of the Nonlinear regression dialog.

2. A drop down menu lets you choose to enter a new equation, clone an existing one, or import an equation from a saved .PZF file. Choose to clone.

3. Edit the duplicated (cloned) copy. You can change the equation, the rules for initial values, constraints and transforms to report.

4. The default name of the cloned equation is the original name followed by a digit. Change to a more informative name.

The cloned equation is an exact duplicate of the equation you started with except for a digit appended to the equation name. Cloning does more than duplicate the math. It also includes the description, hint, rules for initial values, default constraints, and transforms to report.
3.7.4 How to: Manage your list of equations

Deleting user-defined equations

Every time you enter an equation, or edit an equation in a file you are working on, Prism adds that equation to your list of user-defined equations. Over time, this list can get too long to easily work with.

When working with the Fit tab of the nonlinear regression dialog, click on any user-defined equations, then click **Delete** or **Delete All**.

Don't hesitate to delete equations. It won't affect any files that use the equation you deleted. If you open one of these files, and change the parameters of the nonlinear regression, Prism will automatically add the equation back to your list. You can also import an equation from any Prism file onto your list of user-defined equations.

Changing the order of equations

Equations appear on the list in the order they were added. To change the order of equations in the list, select an equation and then click **Move up** or **Move down**. You can only change the order of user-defined equations, not the order of built-in equations.

Renaming an equation

The equation name helps you choose it in the future. It also appears on the analysis results. You are not stuck with the name you originally gave it. To **rename** an equation, select the equation, and then click **Edit**. Change the name and click OK.

Sending an equation to a colleague

The easiest way to send an equation to a colleague is to send a file that uses that equation. Ask your colleague to go to the results page, click the button in the upper left corner of the results table to bring up the
Parameters: Nonlinear regression dialog. Then click OK. The equation will be appended to the list of equations.

### 3.7.5 Syntax of user-defined equations

**General syntax**

- Variable and parameter names must not be longer than 13 characters.

- If you want to use two words to name a variable, separate with the underscore character, for example Half-Life. Don't use a space, hyphen or period.

- Prism does not distinguish between upper and lower case letters in variable, parameter or function names.

- Use an asterisk (*) to indicate multiplication. Prism does not always recognize implied multiplication. To multiply A times B, enter “A*B” and not “AB”.

- Use a caret (^) to indicate power. For example, "A^B" is A to the B power.

- Use parentheses as necessary to show the order of operations. To increase readability, substitute brackets [like this] or braces {like this}. Prism interprets parentheses, brackets, and braces identically. Don’t make any assumptions about the order of precedence. Include enough parentheses so there is no ambiguity about how the equation is evaluated.

- Use a single equals sign to assign a value to a variable.

- You don't need any special punctuation at the end of a statement.

- To enter a long line, type a backslash (\) at the end of the first line, then press Return and continue. Prism treats the two lines as one.

- To enter a comment, type a semicolon (;) and then the text. Comments can begin anywhere on a line.
• You can use many **functions**, most which are similar to those built-in to Excel.

• Be careful not to use the name of a built-in function as a parameter name. For example, since *beta* is the name of a function, you cannot name a parameter *beta*.

**IF-THEN relationships**

Prism allows you to introduce some branching logic through use of the IF function. The syntax is:

\[ \text{IF (conditional expression, value if true, value if false).} \]

You can precede a conditional expression with NOT, and can connect two conditional expressions with AND or OR. Examples of conditional expressions:

- MAX>100
- Ymax=Constraint
- (A<B or A<C)
- NOT(A<B AND A<C)
- FRACTION<>1.0
- X<=A and X>=B

Prism's syntax is that of most computer languages: "<>" means not equal to, "<=" means less than or equal to, and ">=" means greater than or equal to.

Here is an example.

\[ Y= \text{If} (X<X0, \text{Plateau, Plateau} \times \exp(-K \times X)) \]

In this example, if \( X \) is less than \( X0 \), then \( Y \) is set equal to Plateau. Otherwise \( Y \) is computed as \( \text{Plateau} \times \exp(-K \times X) \). This approach is useful for **segmental regression**.

You may also insert a conditional expression anywhere in an equation, apart from an If function. A conditional expression evaluates as 1.0 if true and 0.0 if false. Example:

\[ Y=(X<4) \times 1 \times (X>=4) \times 10 \]
When $X$ is less than 4, this evaluates to $1\times 1 + 0\times 10 = 1$. When $X$ is greater than 4, this evaluates to $0\times 1 + 1\times 10 = 10$.

### 3.7.6 Multiline models

Equations can be written over several lines. Here is an example, the mixed model enzyme inhibition model built-in to Prism:

$$
V_{\text{max App}} = \frac{V_{\text{max}}}{1 + \frac{I}{(\text{Alpha} \times K_i)}} \\
K_{\text{m App}} = \frac{K_m \times (1 + \frac{I}{K_i})}{1 + \frac{I}{(\text{Alpha} \times K_i)}} \\
Y = \frac{V_{\text{max App}} \times X}{(K_{\text{m App}} + X)}
$$

Prism follows the convention of all computer languages. It starts at the top and goes down.

First it computes the intermediate variable $V_{\text{max App}}$. It knows this is an intermediate variable, and not a parameter to fit, because it appears to the left of the equals sign.

Next Prism computes the value of $K_{\text{m App}}$.

Finally it uses those two values to compute $Y$.

Math textbooks tend to write equations in the opposite order. A math text might first define $Y$ as a function of $V_{\text{max App}}$ and $K_{\text{m App}}$, and then lower on the page define how to calculate $V_{\text{max App}}$ and $K_{\text{m App}}$ from $V_{\text{max}}$, $K_m$, Alpha and I. Prism (like all computer languages) requires that you define an intermediate variable before you use it.

Here is a second example:

$$
\text{Specific} = X \times B_{\text{max}} / (X + K_d) \\
\text{Nonspecific} = NS \times X \\
<A> Y = \text{Specific} + \text{Nonspecific} \\
<B> Y = \text{Nonspecific}
$$

The first line calculates the intermediate variable Specific. The second line defines the intermediate variable Nonspecific.
The third line is preceded by <A>. This means that this line only applies to data set A. The second line is preceded by <B> so only applies to data set B. This allows the model to fit a table of data where column A is the total binding and column B is the nonspecific binding. Read more about the syntax used to specify that a particular line only applies only to selected data sets.

You can define constants right in the multiline equation. This makes sense for defining true constants, whose value will never change. If the constant is something like a concentration that changes from experiment to experiment, it is better to not define it in the equation itself, but rather define it in the Constrain tab. If it first appears on the right side of the equation, Prism will treat it like a parameter. You can use the Constrain tab to fix that parameter to a constant value. If the variable name appears first to the left of the equals sign, it is used only within the equation and won't appear in the constraints tab. For example, this line defines Pi:

\[ \pi = 3.141529 \]

### 3.7.7 Limitations when entering equations

**Functions must be \( Y = f(X) \)**

When you enter an equation into Prism, the independent variable must be 'X' and the dependent variable must be 'Y'. So if you measure a voltage as a function of time, you cannot enter an equation that defines \( V \) as a function of \( t \). It must define \( Y \) as a function of \( X \).

**No models with more than one X variable**

Prism does not calculate multiple regression, so cannot fit models with two or more independent (X) variables. But note that you can define a parameter to be a column constant, in which case its value comes from the column titles. In some cases, you can think of these column constants as being a second independent variable.

**Ways to evade this limitation.**

**Model complexity**

Prism compiles your equation into an internal format it uses to calculate the math efficiently. If the compiled version of your equation won't fit in
the space Prism sets aside for this purpose, it reports that the equation is "too complex".

If you see this message, don't give up. You can usually rewrite an equation to make it less complex. Do this by defining an intermediate variable that defines combinations of variables. For example if your equation uses the term "K1+K2" four times, you reduce complexity (but keep exactly the same mathematical meaning) by defining an intermediate variable at the top of your equation (say, K1P2=K1+K2) and then using that intermediate later in the equation. That way Prism has fewer steps to store.

3.7.8 Entering a differential equation

An example. Exponential decay

A single phase exponential decay is defined by this equation.

\[ Y = Y_0 \cdot \exp(-K \cdot X) \]

Simple calculus shows that this has the property that the derivative of \( Y \) is proportional to the value of \( Y \).

\[ \frac{dY}{dX} = -K \cdot Y \]

How to fit a differential equation with Prism.

Prism can fit a model defined by a differential equation.

Choose the differential equation type at the top of the Equation dialog, and define \( Y' \) (the derivative of \( Y \) with respect to \( X \)) as a function of \( X \) and parameters. For the example, enter:

\[ Y' = -K \cdot Y \]

That's it. Putting an apostrophe after the \( Y \) on the left side of the equation tells Prism that you are defining the derivative of \( Y \) with respect to \( X \).
Notes

- Prism does not understand the other nomenclature for differential equations. Don't try to define an equation that starts with "dY/dX = ".

- Note that X doesn't actually appear in the equation. That's ok. It is there in spirit, since Y' defines the derivative of Y with respect to X.

- When you look at that equation, there appears to be only one parameter, K. In fact, the equation has two parameters. Prism generates a parameter Y[0], which is the value of Y at X=X0.

- When you go to add constraints and initial values, Y[0] appears just like the other parameter K.

- You can set X0 to any constant value you want, but it is usually set to 0.0. If you want to choose a different value, set it on the bottom of the initial values tab of the dialog that defines the equation. Note this is the dialog that defines the equation, not the dialog used for each fit.

- Fitting a differential equation requires more calculations, so it takes noticeably more time that fitting the usual kind of equation.

- It is only possible to define Y'. It is not possible to use differential equations to define intermediate variables. This would be useful for fitting compartmental models, but Prism cannot (yet) fit this kind of model.

- With models defined as differential equations, Prism 6 did not allow you to define different models for different data sets using the <A>..<B>.. notation. Prism 7 does allow this. But note that in every case, you must define Y'. It is not possible to define Y' for some data sets and define Y for others.

- Prism 6 always fit the Y[0] value and shared its value for all data sets. Prism 7 lets you set it to a constant value, and lets you share it among data sets or not.

3.7.9 Entering an implicit equation

An implicit equation is one where Y appears on both sides of the equals
sign. Prism 6 will (try to) fit models defined by implicit equations to your data.

**Example of an implicit equation**

The model for total binding at equilibrium to a binding site that follows the law of mass action is

\[
Y = NS*X + Bmax*X/(Kd + X)
\]

That form of the model assumes that \( X \) is the free concentration of ligand. Since \( X \) is actually the concentration you entered, this model assumes that a tiny fraction of the ligand binds so \( X \) is

What if a substantial fraction of the ligand binds? A much more complicated model is built into Prism to handle this situation. But in other similar situations, the model may not have been derived and it may be very difficult or even impossible to derive one. Implicit equations to the rescue.

In the situation above, it is easy to write an implicit equation. There are three places in the model where \( X \) appears. The \( X \) values you enter are the total concentration of ligand. The value that enters into the model must be the free concentration, which is the total concentration minus the concentration that bound. Assuming \( X \) and \( Y \) are both in the same units, we simply replace \( X \) with \( X-Y \) in all three places:

\[
Y = NS*(X-Y) + Bmax*(X-Y)/(Kd + (X-Y))
\]

When you actually go to fit the data, you'll probably want to change the equation a bit (as shown below) to handle unit conversions so the \( Kd \) is reported in nM rather than cpm.

**How to fit implicit equations with Prism**

Choose the implicit equation type at the top of the Equation dialog, and define the equation with \( Y \) on both sides of the equals sign.
That's it. You'll need to define constraints and initial values as with any user-defined equation.

**Notes**

- In this example you don't actually see Y on both sides of the equals sign in the same equation line. But the first line puts Y on the right side of the equations sign, and the fourth line puts it on the right side. That makes the equation implicit.

- If you subtract Y from X, as in the example here, then X and Y must both be entered in the same units. Here both are entered as radioactivity counts per minute (cpm). Of course, it would make no sense to subtract Y from X if Y were in cpm and X were in nM.

- Prism finds it "harder" to fit implicit equations than ordinary ones. You may have to fuss with initial values and constraints to get it to work. The calculations take much longer, although this may not be noticeable with small data sets and fast computers.

- While Y appears on both sides of the equals sign in an explicit equation, X must appear only on the right side of the equals sign.

- In this particular example, the explicit equation has been derived and is even built into Prism. The file you can download (link below) fits the data both ways (explicit equation and implicit equation) and the results are identical. In other cases, it may be difficult or impossible to derive an explicit equation.
### 3.7.10 Available functions for user-defined equations

**Allowed syntax**

<table>
<thead>
<tr>
<th>Function</th>
<th>Explanation</th>
<th>Excel equivalent</th>
</tr>
</thead>
<tbody>
<tr>
<td>abs(k)</td>
<td>Absolute value. If k is negative, multiply by 1.</td>
<td>abs(k)</td>
</tr>
<tr>
<td>arccos(k)</td>
<td>Arccosine. Result is in radians.</td>
<td>acos(k)</td>
</tr>
<tr>
<td>arccosh(k)</td>
<td>Hyperbolic arc cosine.</td>
<td>acosh(k)</td>
</tr>
<tr>
<td>arcsin(k)</td>
<td>Arcsine. Result is in radians.</td>
<td>asin(k)</td>
</tr>
<tr>
<td>arcsinh(k)</td>
<td>Hyperbolic arcsin. Result in radians.</td>
<td>asinh(k)</td>
</tr>
<tr>
<td>arctan(k)</td>
<td>Arctangent. Result is in radians.</td>
<td>atan(k)</td>
</tr>
<tr>
<td>arctanh(k)</td>
<td>Hyperbolic tangent. K is in radians.</td>
<td>atanh(k)</td>
</tr>
<tr>
<td>arctan2(x,y)</td>
<td>Arctangent of y/x. Result is in radians.</td>
<td>atan2(x,y)</td>
</tr>
<tr>
<td>besselj(n,x)</td>
<td>Integer Order J Bessel, N=0,±1, ±2...</td>
<td>besselj(x,n)</td>
</tr>
<tr>
<td>bessely(n,x)</td>
<td>Integer Order Y Bessel, N=0,±1, ±2...</td>
<td>bessely(x,n)</td>
</tr>
<tr>
<td>besseli(n,x)</td>
<td>Integer Order I Modified Bessel, N=0,±1, ±2...</td>
<td>besseli(x,n)</td>
</tr>
<tr>
<td>besselk(n,x)</td>
<td>Integer Order K Modified Bessel, N=0,±1, ±2...</td>
<td>besselk(x,n)</td>
</tr>
<tr>
<td>beta(j,k)</td>
<td>Beta function.</td>
<td>exp(gammaln(j) +gammaln(k) -gammaln(j+k))</td>
</tr>
<tr>
<td>Function</td>
<td>Explanation</td>
<td>Excel equivalent</td>
</tr>
<tr>
<td>------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>-----------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>binomial(k,n,p)</td>
<td>Binomial. Probability of k or more “successes” in n trials, when each trial has a probability p of “success”.</td>
<td>1 - binomdist(k,n,p,true) + binomdist(k,n,p,false)</td>
</tr>
<tr>
<td>chidist(x2,v)</td>
<td>P value for chi square equals x2 with v degrees of freedom.</td>
<td>chidist(x2,v)</td>
</tr>
<tr>
<td>ceil(k)</td>
<td>Nearest integer not smaller than k.</td>
<td>(no equivalent)</td>
</tr>
<tr>
<td>cos(k)</td>
<td>Cosine. K is in radians.</td>
<td>cos(k)</td>
</tr>
<tr>
<td>cosh(k)</td>
<td>Hyperbolic cosine. K is in radians.</td>
<td>cosh(k)</td>
</tr>
<tr>
<td>deg(k)</td>
<td>Converts k radians to degrees.</td>
<td>degrees(k)</td>
</tr>
<tr>
<td>erf(k)</td>
<td>Error function.</td>
<td>2<em>normsdist(k</em>sqrt(2))-1</td>
</tr>
<tr>
<td>erfc(k)</td>
<td>Error function, complement.</td>
<td>2 - 2<em>normsdist(k</em>sqrt(2))</td>
</tr>
<tr>
<td>exp(k)</td>
<td>e to the kth power.</td>
<td>exp(k)</td>
</tr>
<tr>
<td>floor(k)</td>
<td>Next integer below k.</td>
<td>(no equivalent)</td>
</tr>
<tr>
<td>fdist(f,v1,v2)</td>
<td>P value for F distribution with v1 degrees of freedom in the numerator and v2 in the denominator.</td>
<td>fdist(f,v1,v2)</td>
</tr>
<tr>
<td>gamma(k)</td>
<td>Gamma function.</td>
<td>exp(gammaln(k))</td>
</tr>
<tr>
<td>gammaln(k)</td>
<td>Natural log of gamma function.</td>
<td>gammaln(k)</td>
</tr>
<tr>
<td><strong>Function</strong></td>
<td><strong>Explanation</strong></td>
<td><strong>Excel equivalent</strong></td>
</tr>
<tr>
<td>-------------------</td>
<td>------------------------------------------</td>
<td>---------------------------------------</td>
</tr>
<tr>
<td>hypgeometricm(a,b,x)</td>
<td>Hypergeometric M.</td>
<td>(no equivalent)</td>
</tr>
<tr>
<td>hypgeometricu(a,b,x)</td>
<td>Hypergeometric U.</td>
<td>(no equivalent)</td>
</tr>
<tr>
<td>hypgeometricf(a,b,c,x)</td>
<td>Hypergeometric F.</td>
<td>(no equivalent)</td>
</tr>
<tr>
<td>ibeta(j,k,m)</td>
<td>Incomplete beta.</td>
<td>(no equivalent)</td>
</tr>
<tr>
<td>if(condition, j, k)</td>
<td>If the condition is true, then the result is j. Otherwise the result is k. See details below.</td>
<td>(similar in excel)</td>
</tr>
<tr>
<td>igamma(j,k)</td>
<td>Incomplete gamma.</td>
<td>gammadist(k, j, 1,TRUE)</td>
</tr>
<tr>
<td>igammac(j,k)</td>
<td>Incomplete gamma, complement.</td>
<td>1 - gammadist(k, j, 1,TRUE)</td>
</tr>
<tr>
<td>int(k)</td>
<td>Truncate fraction.</td>
<td>trunc()</td>
</tr>
<tr>
<td>ln(k)</td>
<td>Natural logarithm.</td>
<td>ln(k)</td>
</tr>
<tr>
<td>log(k)</td>
<td>Log base 10.</td>
<td>log10(k)</td>
</tr>
<tr>
<td>max(j,k)</td>
<td>Maximum of two values.</td>
<td>max(j,k)</td>
</tr>
<tr>
<td>min(j,k)</td>
<td>Minimum of two values.</td>
<td>min(j,k)</td>
</tr>
<tr>
<td>j mod k</td>
<td>The remainder (modulus) after dividing j by k.</td>
<td>mod(j,k)</td>
</tr>
<tr>
<td>psi(k)</td>
<td>Psi (digamma) function. Derivative of the gamma function.</td>
<td>(no equivalent)</td>
</tr>
<tr>
<td>Function</td>
<td>Explanation</td>
<td>Excel equivalent</td>
</tr>
<tr>
<td>----------</td>
<td>-------------</td>
<td>------------------</td>
</tr>
<tr>
<td>rad(k)</td>
<td>Converts k degrees to radians.</td>
<td>radians(k)</td>
</tr>
<tr>
<td>sgn(k)</td>
<td>Sign of k.</td>
<td>sign(k)</td>
</tr>
<tr>
<td></td>
<td>If k&gt;0, sgn(k)=1.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>If k&lt;0, sgn(k)= -1.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>If k=0, sgn(k)=0.</td>
<td></td>
</tr>
<tr>
<td>sin(k)</td>
<td>Sine. K is in radians.</td>
<td>sin(k)</td>
</tr>
<tr>
<td>sinh(k)</td>
<td>Hyperbolic sine. K is in radians.</td>
<td>sinh(k)</td>
</tr>
<tr>
<td>sqr(k)</td>
<td>Square.</td>
<td>k*k</td>
</tr>
<tr>
<td>sqrt(k)</td>
<td>Square root.</td>
<td>sqrt(k)</td>
</tr>
<tr>
<td>tan(k)</td>
<td>Tangent. K is in radians.</td>
<td>tan(k)</td>
</tr>
<tr>
<td>tanh(k)</td>
<td>Hyperbolic tangent. K is n radians.</td>
<td>tanh(k)</td>
</tr>
<tr>
<td>tdist(t,v)</td>
<td>P value (one-tailed) corresponding to specified value of t with v degrees of freedom. T distribution.</td>
<td>tdist(t,v,1)</td>
</tr>
<tr>
<td>zdist(z)</td>
<td>P value (one-tailed) corresponding to specified value of z. Gaussian distribution.</td>
<td>normsdist(z)</td>
</tr>
</tbody>
</table>

### 3.7.11 Fitting different segments of the data to different models

**What is segmental regression?**

In some situations you may wish to fit different models to different portions of your data. This is called **segmental regression**. This often occurs in kinetic experiments where you add a drug or perform some sort
of intervention while recording data. The values collected before the intervention follow a different model than those collected afterwards.

If you didn't perform an intervention at a particular time along the X axis, it is unlikely that this kind of segmental regression is the analysis of choice.

**How to perform segmental regression with Prism**

Although Prism has no built-in way to fit different equations to different portions of the data, you can achieve that effect using a user-defined equation containing the IF function.

In this example, you collected data that established a baseline early in the experiment, up to "Start". You then added a drug, and followed the outcome (Y) as it increased towards a plateau. Prior to the injection, the data followed a horizontal line; after the injection the data formed an exponential association curve.

\[
Y_1 = \text{BASELINE} \\
Y_2 = \text{BASELINE} + \text{SPAN} \times (1 - \exp(-K \times (X - \text{START}))) \\
Y = \text{IF}[(X < \text{START}), Y_1, Y_2]
\]

It is easiest to understand this equation by reading the bottom line first. For X values less than START, Y equals Y1, which is the baseline. Otherwise, Y equals Y2, which is defined by the exponential association equation.

This equation has two intermediate variables (Y1 and Y2). Prism can fit the four true variables: START, SPAN, K, and BASELINE.
In many cases, you will make START a constant equal to the time of the experimental intervention. If you want Prism to fit START, choose an initial value carefully.

This kind of model is most appropriate when X is time, and something happens at a particular time point to change the model. In the example above, a drug was injected at time=Start.

**How to fit a model to only a portion of the data**

The Range tab of the nonlinear regression dialog lets you define an X range that determines which points are fit and which are ignored.

### 3.7.12 Fitting different models to different data sets

**Writing a model so some lines only apply to selected data sets**

Prism can only fit data from one table at a time, and all the data sets must be fit by a single equation you select or create. But you can create this equation in such a way that different data sets are fit to mathematically distinct equations.

When entering a user defined equation, you use multiple lines to define intermediate variables. Each line can have a prefix that specifies which data sets that line will be used for:

<table>
<thead>
<tr>
<th>Prefix</th>
<th>This line applies to</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;C&gt;</td>
<td>Data set C only</td>
</tr>
<tr>
<td>~B</td>
<td>All data sets, except B.</td>
</tr>
<tr>
<td>A:D</td>
<td>Data sets A-D.</td>
</tr>
<tr>
<td>~A:D</td>
<td>All data sets except A-D.</td>
</tr>
<tr>
<td>A:J,3</td>
<td>Data sets A,D,G, and J (every third data set between A and J.)</td>
</tr>
<tr>
<td>~A:J,3</td>
<td>All data sets except A,D,G, and J.</td>
</tr>
</tbody>
</table>
Here is an example. It fits column A to a model that defines total binding and column B to a model that defines nonspecific binding only. The first two lines of the equation are evaluated for all data sets, the third line is only evaluated for data set A, while the last line is only evaluated for data set B. To fit this model, you would want to set the constraint that the parameter NS is shared between data sets.

\[
\text{Specific} = X \times B_{\text{max}} / (X + K_d) \\
\text{Nonspecific} = \text{NS} \times X \\
\text{A} = \text{Specific} + \text{Nonspecific} \\
\text{B} = \text{Nonspecific}
\]

**Ambiguity when you select noncontinuous range of data sets to analyze**

\[
\text{A} = 1 / (1 + K_a \times X^h) \\
\text{C} = (K_a \times X^h) / (1 + K_a \times X^h)
\]

The equation above seems clear at first. The first line is for data set A, and the second line is one for data set C. But what if you asked Prism to only analyze data sets A and C, skipping B? Now it is a bit ambiguous. Does \(\text{C}\) mean data set C, or does it mean the third data set included in the analysis? Prism uses the second definition. So you need to enter the equation like this.

\[
\text{A} = 1 / (1 + K_a \times X^h) \\
\text{B} = (K_a \times X^h) / (1 + K_a \times X^h)
\]

Now the second line is for the second data set, which in this example is data set C (since only A and C were selected in the Analyze dialog).

### 3.7.13 Column constants

**What is a column constant?**

When you fit a number of datasets at once, you can use the column title as a second independent variable. We call this constraining a parameter to be a column constant. This is best seen by example.

**How to enter column constants**

To see how column constants work, create a new XY table using the sample data file: Enzyme kinetics - Competitive inhibition.

The data table has one X column, and four Y columns, each representing a different concentration of inhibitor. The inhibitor concentrations are entered as column title.
Note that Prism reads only the number in the column title. In this example, the units are specified as micromolar, but Prism ignores this and simply reads the numbers. It does not do any unit conversion.

**Specifying a column constant when fitting data**

When choosing an initial value, choose from the drop down list either "Mean of column title value" or "log(Mean of column title value)."

```
<table>
<thead>
<tr>
<th>Initial Value</th>
<th>Rule</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>*(Mean of column title values)</td>
</tr>
</tbody>
</table>
```

**Example**

To fit the sample data above, click Analyze, choose nonlinear regression, choose the Enzyme Kinetics panel of equations and choose Competitive enzyme kinetics. The equation is built in, but if you click the Details button you can see the math.

\[
K_{\text{mobs}} = K_m (1 + [I] / K_i) \\
Y = V_{\text{max}} X / (K_{\text{mobs}} + X)
\]

The first line defines an intermediate variable (\(K_{\text{mobs}}\), the observed Michaelis-Menten constant in the presence of a competitive inhibitor), which is a function of the Michaelis-Menten constant of the enzyme (\(K_m\)), the concentration of inhibitor (\([I]\)), and the competitive inhibition constant (\(K_i\)).
The second line computes enzyme velocity (Y) as a function of substrate concentration (X) and KMapp.

This model is defined with I constrained to being a data set constant, which means its value comes from the column titles. In this example, therefore, I=0 when fitting column A, I=5 when fitting column B, etc. The 'μM' in the title is ignored by Prism -- it doesn't do any unit conversions.

The other three parameters (Km, Ki and Vmax) are defined to be shared, so Prism fits one best-fit value that applies to the entire family of datasets.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>KM</td>
<td>Shared value for all data sets</td>
</tr>
<tr>
<td>I</td>
<td>Data set constant (=column title)</td>
</tr>
<tr>
<td>Ki</td>
<td>Shared value for all data sets</td>
</tr>
<tr>
<td>VMAX</td>
<td>Shared value for all data sets</td>
</tr>
</tbody>
</table>

Prism determined the maximum velocity of the enzyme with no inhibitor (Vmax in the same units as the Y values you entered), the Michaelis-Menten constant of the enzyme with no inhibitor (Km, in the units used for X values) and the competitive inhibition constant (Ki, in units used for the column constants). Note that I is not a parameter to be fit, but rather takes on constant values you entered into the column titles. KmObs is not a parameter to be fit, but is rather an intermediate variable used to define the model.

Learn more about competitive enzyme inhibition.

Summary. The advantage of column constants

By using column constants and global fitting (shared parameters), this example determined a parameter (Ki) whose value cannot be determined from any one dataset, but can only be determined by examining the relationships between datasets.

3.7.14 Defining equation with two (or more) independent variables

Prism is designed to perform nonlinear regression with one independent (X) variable. But, with a bit of cleverness, it is possible to also fit data with two independent variables. There are three separate cases:
Fitting a family of curves

The graph above shows the sample data for competitive enzyme kinetics, and shows how to fit a family of curves. Here, each curve shows enzyme activity as a function of substrate concentration. The curves differ by the presence of various amounts of an inhibitor. There are, essentially, two independent variables: substrate concentration, and inhibitor concentration.

The substrate concentration is entered into the X column and the inhibitor concentration into the column titles.

When fitting the data, the Constrain tab of the nonlinear regression dialog is used to define the parameter I as dataset constant whose value comes from the column titles (so is not fit by regression), and to share the values of all the other parameters so there is one global fit of all the data.
Fitting data where each value has two independent variables

But what if each value you collect has two independent variables? You don't have a family of curves. Rather you have one set of Y values, each associated with two X variables. Follow these steps to enter the data:

1. Create an XY data table, with whatever form of subcolumns fits your data.

2. Enter the first Y value in row 1 of column A.

3. Enter the first independent variable corresponding to that Y into first row of the X column.

4. Enter the other independent variable for that first Y value as the column title of column A.

5. Enter the second Y value in row 2 of column B.

6. Enter the first independent variable for that Y into the second row of the X column.

7. Enter the other independent variable for that Y as the column title of column B.
8. Continue with the rest of the data. You'll be entering data diagonally down the table. Since Prism can only have 104 columns, you will be limited to 104 data points, each with a Y value and two X values.

9. When you are done, you'll have the same number of rows as Y columns.

```
<table>
<thead>
<tr>
<th>X</th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
<th>E</th>
<th>F</th>
<th>G</th>
</tr>
</thead>
<tbody>
<tr>
<td>X Title</td>
<td>0.006556</td>
<td>0.000542</td>
<td>0.000529</td>
<td>0.000505</td>
<td>0.000483</td>
<td>0.000462</td>
<td>0.000444</td>
</tr>
<tr>
<td>X</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
<td>Y</td>
</tr>
<tr>
<td>0.000000</td>
<td>0.000000</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.117000</td>
<td></td>
<td>0.015103</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.229000</td>
<td></td>
<td></td>
<td>0.022470</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.436000</td>
<td></td>
<td></td>
<td></td>
<td>0.035167</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.628000</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.041316</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.730000</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.046776</td>
<td></td>
</tr>
<tr>
<td>0.958000</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.046327</td>
</tr>
</tbody>
</table>
```

When you fit the data, keep in mind the following:

- "X" refers to the values you enter into the X column. Use some other name ("X2" or whatever you want) for the other independent variable.

- In the constraints tab, define that parameter to be a data set constant equal to the column title. You don't want Prism to fit that parameter, but rather to get its value from the column titles.

- For all other parameters in the model, use the Constraint tab to share the value among data sets. Prism sees each column as a data set. In this data table, each data set has only one value so Prism can't possibly fit anything unless all the parameters are shared.

Prism will report the nonlinear regression results clearly. All the results you want will be in the final "global" column. Prism cannot graph the results since each data set has only one point, and Prism cannot plot 3D graphs.

**Three independent variables**

Prism is not really designed to fit models with three independent variables, but you can get it to do so in some circumstances.
Enter your data as above, with one independent variable as X and the second as column titles. Make sure each column of data has only one value for the third independent variable. Then define that third variable in a user-defined equation. Let’s imagine that the third variable has the name Z. Add syntax like this to your equation:

\(<A>\ Z=2.34\)
\(<B>\ Z=45.34\)
\(<C>\ Z=\ -23.4\)
\(<D>\ Z=12.45\)

That tells Prism to assign Z one value for column A, a different value for column B, etc. This approach gives you three independent variables, with some constraints:

- The X column is one independent variable, with one value for each row.
- The column titles form a second independent variable with one value per data set column.
- The Z values defined within the equation for a third independent variable, also with one value per data set.
3.7.15 Reparameterizing an equation

Reparameterizing an equation does not change the best-fit curve

You want to fit the sigmoidal enzyme kinetics data to a standard model. But there are two forms of that model that are commonly used:

\[ Y = \frac{V_{\text{max}} \times X^h}{(K_{\text{half}}^h + X^h)} \]
\[ Y = \frac{V_{\text{max}} \times X^h}{(K_{\prime} + X^h)} \]

The two are equivalent, with \( K_{\prime} \) equal to \( K_{\text{half}}^h \), so the two fits will generate exactly the same curve, with the same sum-of-squares, the same \( R^2 \), and the same numbers of degrees of freedom. Even though the two equations express the same model, they are written differently. The fancy term is that they are parameterized differently.

They both fit \( V_{\text{max}} \) (the maximum activity extrapolated to very high concentrations of substrate) and \( h \) (Hill slope, describing the steepness of the curve). But one model fits \( K_{\text{half}} \) (the concentration needed to obtain a velocity half of maximal) and the other fits \( K_{\prime} \) (a more abstract measure of substrate action).

Which model is preferred?

One way to choose between the two models is to match other text books and papers, so your results can easily be compared to others. Another approach is to choose the form that fits the way you think. For this example, if you prefer to think graphically, choose the \( K_{\text{half}} \). If you think mechanistically, choose \( K_{\prime} \).
But the choice can be more than a matter of convenience and convention. The choice of model can determine the accuracy of the confidence intervals. Read on to see why.

**Distribution of parameters are not always symmetrical**

*Simulations* can determine parameter symmetry. I simulated sigmoidal enzyme kinetics using \( \text{Vmax}=100, \ h=5, \ K_{\text{prime}}=25, \) and Gaussian scatter with a SD equal to 7.5. The X values matched those in the figure above, with triplicate Y values at each X. Prism can do such simulations easily. Using Prism's *Monte Carlo analysis*, I repeated the simulations 5000 times, fit each curve to both forms of the models, and tabulated the best-fit values of \( K_{\text{prime}} \) and \( K_{\text{half}} \), and computed the skewness of each.

![Frequency distribution of \( K_{\text{half}} \) and \( K_{\text{prime}} \)](image)

The distribution of \( K_{\text{half}} \) is quite symmetrical and looks Gaussian. Accordingly, its skewness is close to zero. In contrast, the distribution of \( K_{\text{prime}} \) is quite skewed. Note that a few of the simulation data sets had best-fit values of \( K_{\text{prime}} \) greater than 100. The skewness value (4.89) confirms what is obvious by inspection -- the distribution is far from symmetrical.

**Coverage of the symmetrical asymptotic confidence intervals**

Prior versions of GraphPad Prism, like almost all nonlinear regression programs, computes the confidence interval of a best-fit parameters so the interval is symmetrical around the best fit values. If the uncertainty of a parameter is truly symmetrical, then these confidence intervals can
be interpreted at face value. If the uncertainty is not symmetrical, then the confidence interval will not be accurate.

Ideally, a confidence interval is easy to interpret. A 95% CI has a 95% chance of including the true value of the parameter, and a 5% chance of missing it. When analyzing real data, we never know the value of the true parameter, so never know if the interval includes it or not. But when you simulate data, you know the true values of the parameters, so can quantify the coverage of the confidence intervals. I set up the same simulations mentioned above, fit each data set to both equations, and tabulated whether each confidence interval included the true parameter value or not. This table shows the fraction of 5,000 simulations where the confidence interval did not include the true parameter value (25 for Kprime, and 1.9037 for Khalf).

<table>
<thead>
<tr>
<th></th>
<th>&quot;95% CI&quot;</th>
<th>&quot;99% CI&quot;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ideal</td>
<td>5.0%</td>
<td>1.0%</td>
</tr>
<tr>
<td>Kprime</td>
<td>8.8%</td>
<td>4.8%</td>
</tr>
<tr>
<td>Khalf</td>
<td>5.1%</td>
<td>1.0%</td>
</tr>
</tbody>
</table>

These results show that Khalf is well behaved, as expected given its symmetry (see above). The 95% confidence interval is expected to miss the true value in 5.0% of the simulations. In fact, it happened 5.1% of the time. Similarly, the 99% CI is expected to miss the true value in 1.0% of the simulations, which is exactly what happened. In contrast, Kprime is less well behaved. The intervals computed to be 95% confidence intervals were not wide enough so missed the true value in 8.8% of the simulations. The 99% intervals were similarly not wide enough so missed the true value in 4.8% of the simulations. Thus the confidence intervals computed to be 99% intervals, actually turned out to be 95% intervals.

These simulations show the advantage of choosing the equation that fits Khalf, rather than the one that fits Kprime. Khalf has a symmetrical distribution so the confidence intervals computed from these fits can be interpreted at face value. In contrast, Kprime has an asymmetrical distribution and its confidence intervals cannot be interpreted at face value.
Coverage of the asymmetrical likelihood confidence intervals

A new feature in Prism 7 is that it can compute asymmetrical likelihood confidence intervals. These intervals will have the same coverage no matter how the equation is parameterized. The 95% CI of Khalf failed to include the true value (25) in only 5.3% of 10,000 simulations. When the equation was parameterized to fit Kprime, the 95% CI failed to include the true value (1.9037) in 5.1% of 10,000 simulations.

This demonstrates the advantage of using asymmetrical confidence intervals. You'll get meaningful confidence intervals no matter how you parameterize the equation. You can choose the parameterization that fits parameters that match the way you think about the system or are easy to explain. You don't need to choose a parameterization that leads to a symmetrical distribution of parameters.

Hougaard's skewness

The results above were obtained by running numerous simulations. There is an easier way to figure out how symmetrical a parameter is. Hougaard's skewness (new in Prism 6) quantifies the asymmetry of each parameter, computed from the equation, the number of data points, the spacing of the X values, and the values of the parameters.

For the simulated data set, Hougaard's skewness is 0.09 for Khalf and 1.83 for Kprime. A rule of thumb is to expect problems from asymmetry when the absolute value of the Hougaard's skewness is greater than 0.25, and big problems when the value is greater than 1.0. So Hougaard's skewness tells you that the confidence intervals will be accurate when you fit Khalf, but not be so accurate when you fit Kprime.

Note that Hougaard's skewness can be reported as part of the results of nonlinear regression (choose in the Diagnostics tab). No simulations are required.

Bottom line

Models can often be parameterized in multiple ways. You'll get the same curve either way, but choosing an optimum parameterization ensures that the confidence intervals for the parameters are believable. The best way to assess various parameterizations is to ask Prism to report the value of Hougaard's measure of skewness for every parameter.
Simulations take a bit more work, but let you see how symmetrical a parameter is.

File used for this example  Asymmetrical CI file.

### 3.7.16 Rules for initial values

#### Why initial values?

Before it can perform nonlinear regression, Prism must have initial (estimated) values for each parameter in the equation. It then iteratively modifies these values to improve the fit of the model to the data. The initial values must be reasonable, so that the curve defined by those initial values has generally the shape of the data, and doesn't go awfully far from the data points. If the initial values are really bad, Prism may not be able to fit the model.

The need for initial values is not unique to Prism, but is intrinsic to nonlinear regression.

#### Why rules for initial values?

Prism can store rules (algorithms) to calculate initial values from the range of your data. If you choose a built-in equation, the rules are taken care of. If you enter your own (user-defined) equation, you can also enter rules to define the initial values of each parameter. If you enter a good set of rules, then your equation will work in the future with no need to think about initial values.

If you don't enter rules for initial values, you will need to enter the initial values for every variable, for every data set, every time you fit data.

If you enter a poor set of rules, then the initial values will not generate a curve that goes near the data points, and it is likely that Prism will not be able to fit the model to the data.

#### How to enter a rule

While entering or editing a user-defined equation, click on the tab labeled "Rules for initial values". For each parameter, enter a number in the first column and select a value to multiply or divide by.
Most often, you'll use the choices based on YMIN, YMID or YMAX, the minimum, middle (mean of minimum and maximum), and maximum Y value in your data set, or the corresponding XMIN, XMID or XMAX.

You can also ask Prism to interpolate, for example to base the initial value on the interpolated value of X at the value of Y half way between the smallest and largest value (XatYMID) and related values.

**Initial value, to be fit**

The first choice on the drop-down list is "(Initial value, to be fit)". This means that the value you entered in the box to the left will be the initial value for all data sets. The initial value will not depend on the range of the data (it will not be calculated); it will be the actual value you entered.

**Initial value based on the range of the data**

Most of the rules are based on the range of the X and Y values.

The abbreviation YMIN is the minimum value of Y; YMAX is the maximum value, and YMID is the average of YMIN and YMAX. For example, if you enter "0.5" in the first column and select "YMAX" in the second column, Prism sets the initial value to half of YMAX (which differs for each data set).

The phrase "X at YMID" refers to the X value that corresponds to the mean of the highest and lowest Y values.  [Calculation details.](#)
**SIGN(YatXmax - YatXmin)**

It equals +1 if the curve generally goes up as it goes from left to right:

It equals -1 if the curve generally goes down as it goes from left to right:

It is used as the initial value for the Hill Slope in dose response curves.

Here is an explanation of the math. The `SIGN()` function equals +1 when given a positive number, and -1 when given a negative number. YatXMax is the Y value at the largest X values. YatXmin is the Y value at the smallest X value. (YatXmax - YatXmin) will be positive when the curve goes up, and negative when the curve goes down.

**Mean of column title values (or the log of that mean)**

With some kinds of data, it makes sense to label the data sets numerically. Each column titles has a value.

With these kinds of data, you can access the column title values in two ways:

- Use the Constrain tab and choose "Data set constant (=column title)". That parameter in the model is now a constant, but the value of that constant will differ for each data set column, and gets it value from the column title.

- On the Rules for initial values tab in the dialog that defines an equation, set the initial value of a parameter to equal the mean of those column titles. Or to the logarithm of that mean.

The best way to understand this is with an example. Bring up the sample XY data for competitive enzyme inhibition, and follow the instructions, and inspect the dialog settings.

**Initial, middle and final slope**

New to Prism 7: Base the initial value on the initial, middle, or final slope (dY/dX at the smallest, middle or largest value of X) or its reciprocal.

**Example**

\[ Y = \frac{V_{max} \times X}{(K_m + X)} \]
The equation above is the Michaelis-Menten enzyme kinetics model built in to Prism. If you pick this equation as a built-in equation, the rules for initial values are provided automatically. But if you were entering this as a new user-defined equation, it would be easy to come up with rules.

Vmax is the Y value (enzyme velocity) at very high values of X (substrate). So a reasonable initial value would be to set Vmax equal to the largest Y value in the data set. To do this, enter the value 1.0 and choose the multiplier *Ymax. This means Prism will compute the initial value by multiplying the largest Y value by 1.0. You could argue that Vmax is usually higher than the largest Y value, so should compute the initial value by multiplying 1.25 times Ymax. There are often multiple rules one could justify. These rules only compute the initial value, so there is a lot of room for variation.

Km is the X value (substrate concentration) corresponding to a Y value (enzyme velocity) that is half Vmax. Compute its initial value by multiplying 1.0 times "Value of X at Ymid". Prism then averages the largest and smallest Y values to compute Ymid, and then interpolates from straight connecting lines to find the corresponding X value.
Slope at Yleft, Ymid or Yright

These are the slopes (change in Y divided by change of X) at the left, middle or right of the graph. If you have 30 or more data points, Prism computes the slope from the first, middle or last 10% of the points. If you have fewer than 30 points, Prism computes the slope from the first, middle, or last two points.

If you have more than 30 points, the slope is determined by linear regression on the first, middle or last 10% of the points, rounded down. If there are fewer than 30 points, the slope is determined by the first or last two points. For slopeAtYmid with fewer than 30 points, Prism uses two middle points if there are an even number of points, and otherwise uses the three middle points.

What happens if you don't enter any rules?

If you don't enter rules for every parameter, Prism will insist that you enter initial values each time you fit data to the equation.

How to override the initial values defined by rule

The rules are stored with each equation, and the rules can be edited as part of the dialog that defines and edits equations.

The values defined by the rules are shown in the Initial Value tab of the nonlinear regression dialog. To override the initial values:

1. Select one or several data sets on top of the dialog. Or click "Select all" to select all data sets.

2. Uncheck the "Choose automatically" option.

3. Enter the initial value (as a value, a number, not a rule).

To check how good the initial values are, go to the Diagnostics tab and check the option:

(*) Don't fit the curve. Instead plot the curve defined by the initial values of the parameters.
If the resulting curve doesn't go near the data, revise the initial values (or the rules that generate those values) before asking Prism to fit the model to the data.

### 3.7.17 Default constraints

#### What is a constraint?

Constraining parameters is often essential to getting useful results. Constraints can be used to fix parameters to constant values, to share parameters among data sets (global fitting), and to define one parameter to be a column constant (whose value comes from the column titles in the data table).

#### Defining constraints when defining an equation

You can define default constraints as part of the equation definition. This is useful for constraints that will apply every time the equation is used.

Constraints can be for one parameter (Kfast must be greater than zero) or for the relationship between two parameters (Kfast must be greater than Kslow). But note that you cannot invoke a constraint between two parameters if both parameters also are themselves constrained. In a two phase exponential equation, you may want to constrain both parameters to be greater than zero, and also define one rate constant to be larger than the other (Kfast > Kslow). Prism won't let you do that. What you have to do is define one constraint that Kfast is greater than zero, and another that Kfast is greater than Kslow. But don't put in the constraint that Kslow is greater than zero. That is implied by being larger than Kfast.

#### Defining experimental constants

In some cases, one of the parameters in the equation is an experimental constant. It needs to be set to a constant value, but that value differs each time you run the experiment. Do that by going to the constraints tab of the nonlinear regression dialog every time that equation is used.

When you define the equation, you can add a constraint that reminds anyone using the equation to set that constant value. In the constraints tab when defining the equation, choose the constraint "Constant equal to" but leave the value (to the right) blank. With an equation defined this
way, Prism won’t fit data to that equation until the parameter is defined (or the constraint changed).

3.7.18 Reporting transforms of parameters

When defining an equation, you can also ask Prism to report transforms of the best-fit values on the "Transforms to report" tab.

Prism only reports results when the results of the transform is monotonic. Otherwise the results will be blank. This is true for both symmetric and asymmetric confidence intervals.

Transforms of one parameter

Entering the transform

Enter simple transforms of one parameter. On the left side enter the name you want to give the results. On the right side enter the transform.
**Example**: You fit data to an equation that includes a parameter logEC50 that is the logarithm of an EC50, but you want to also report the EC50. Enter the label 'EC50' on the left and '10^logEC50' on the right (without the quotes).

**Example**: You fit data to an equation that reports a rate constant K, but also want to report the half-life. Enter the label 'HalfLife' on the left and 'ln(2)/K' on the right (without the quotes).

### Confidence intervals - Symmetrical vs. asymmetrical

When transforming a single parameter, choose between symmetrical and asymmetrical confidence intervals via a drop down to the right of the transform.

If you choose to report a symmetrical confidence interval, Prism also reports the SE of that parameter, and both SE and CI are exactly the same as they would have been if you had fit that transformed parameter directly. For example if you fit an exponential decay model to determine the rate constant, and transform that rate constant (take its reciprocal) to compute the time constant, the SE and CI of the time constant will be exactly the same as they would have been had you fit the time constant directly.

If you choose to report an asymmetrical confidence interval, Prism transforms both confidence limits to create the confidence interval of the transformed parameter. In most cases, this transformed confidence interval will not be symmetrical.

Choosing between the two is often not straightforward. The symmetrical intervals give you the results you would have had if you had chosen to express the model differently. The asymmetrical results simply express the same confidence interval on a different scale. You want to compute the confidence intervals on the scale where the true uncertainty is most symmetrical. You can then transform those confidence limits to come up with asymmetrical confidence interval on a different scale.

Prism 4 (and earlier) always reported the asymmetrical intervals for EC50 and half lives, which were automatically calculated for built-in equations, and did not offer the choice of transforming parameters in user-defined models.
If your transform simply changes units, then Prism still offers the two choices, but the choice doesn't matter as both end up being exactly the same. This is the case whenever the transform of parameter K is of the form $a*K+b$

**Interpolating transforms**

**How to interpolate points off the curve**

You can also use these 'transforms' to report values from the curve. The interpolated value and its confidence interval will appear in the results, the same as other transformed parameters.

Use this syntax:

- **Y[value or parameter]**
  The Y value of the curve when X is the value you enter within the brackets. Within the brackets, enter either a number or the name of a parameter and Prism will use the best-fit value of that parameter. You cannot enter a mathematical expression. The Y value will be computed for any X, but confidence intervals will be calculated only when the X value is within the range of the X axis.

- **X[value or parameter]**
  The X value of the curve when Y is the value you enter within the brackets. Within the brackets, enter either a number or the name of a parameter and Prism will use the best-fit value of that parameter. You cannot enter a mathematical expression. Prism searches for the Y value you entered within the range the curve is plotted (Range tab) and extending in each direction a distance equal to half that range. It reports the smallest X value it finds within that range that corresponds to the Y value you entered, and doesn't alert you when the curve oscillates so there are several X values at a particular Y value. If both X and Y are within the axis range, a confidence interval is also calculated.
Example: You fit data to a log(dose) response curve and want to report the antilog of the X value (dose) when Y=50 (which is not always the same as the EC50). For the second example, you would enter "Dose at Y=50' on the left, and '10^X[50]' on the right.

Confidence intervals

The confidence interval for interpolating transforms is computed by interpolation off the confidence bands of the regression curve. You don't have a choice of symmetrical vs. asymmetrical intervals.

Combining two parameters

Usefulness of combining parameters

The bottom half of the tab lets you define combinations of parameters to report. For example, you can report the sum of two parameters, or the ratio. Enter the name you want to give to the transform on the left, and then choose the calculation from the drop-down list (say "P1/P2" for a ratio). Then choose which parameter is P1 and which is P2 in the drop downs on the right.

Confidence intervals

When Prism combines two parameters to come up with a calculated value, it is smart about propagating the errors. The standard error and confidence interval it reports for the calculated variable are exactly the same as they would have been had you rearranged the equation to directly fit that calculated value.

3.8 Plotting a function

Graphing a family of theoretical curves, often called plotting a function, is useful in three contexts:

- To understand the properties of a function. After looking at the graph, go back and change the parameters to see how the graph changes. Or plot a family of curves, where one parameter varies from curve to curve.

- To create graphs for teaching theory.
To understand what initial values would make sense when fitting a model to data with nonlinear regression.

3.8.1 How to: Plot a function

Prism offers a *Plot a function* analysis, but it doesn't in fact analyze any data. Rather it generates curves from an equation you choose and parameters you enter.

**How to: Plot a function**

1. Start from any data table or graph, click Analyze, open the **Generate Curve** folder, and then select **Plot a function**.

2. On the first tab (Function), choose the equation, the starting and ending values of \( X \), and the number of curves you want to plot.

3. On the second tab (Options), choose whether you also want to plot the first derivative, second derivative or integral of the function. The "curve" is actually a set of \( X \) and \( Y \) coordinates that define a series of points that are connected to form the curve. You can choose the number of line segments that will define the curve. There is little reason to change the default (150), unless you want to plot only a portion of the curve on some graphs, in which case you should increase that value.

4. On the third tab (Parameter values), enter the parameter values (or click the fish hook icon to hook analysis or info constants).

**Tips for plotting a function**

**Plotting a family of curves**

If you choose to plot more than one curve (a choice on the first tab), the rest of the dialog works a bit differently.

On the bottom of the second (Options) tab, choose if you want to label each curve manually (enter the labels the top of the third tab) or using the value of one of the parameters.

The top of the third tab lists all the curves you will generate. Select one or more of these curves (or click "select all") and then enter the parameter values below. Often you'll want to first click "select all" and
enter most of the parameters. Then click on one curve at a time, and enter the value for the parameter that varies between curves.

**Plotting each curve on a separate graph**

By default, Prism will make graph that contains all of the curves on one graph. If you want each curve to be on its own graph, go to the results table, click New and choose Graph of existing data. On the dialog that pops up, choose to make one graph for each data set (in this context, a data set is a curve).

**If you plan to zoom in and plot only part of the curve**

The curve is defined, by default, as 150 line segments. That creates a smooth looking curve. But if you then change the range of X values shown on the graph, only a fraction of those line segments will be visible, and the curve may seem coarse. To fix this problem, go back to the parameters dialog, to the Options tab, and increase the number of line segments to a much larger value.

**Combining two curves on one graph**

The graph below combines two plotted functions on one graph. The first time I plotted a function, I chose a Gaussian distribution, with the X ranging from -3 to 3. I set the mean to 0.0, the SD to 1.0, and the amplitude to 100.0 (arbitrary, since I hid that axis). I then repeated that analysis, but this time set the X range from 1.3 to 3.0. I put both curves on one graph (Change.. Add data sets -- remember that a curve generated by this analysis is a "data set" to Prism). For the shorter curve, I chose to create an area fill.
3.8.2 Plotting t, z, F or chi-square distributions

GraphPad Prism can generate probability distributions. This demonstrates Prism's ability to plot functions from user-defined functions, and also the use of hooking info constants to analyses.

Download this Prism file to generate and plot the graphs shown below.
In each case, the simulation generates two (or three) data sets. The first (A) data set plots the entire curve. The equation is written so the second curve (data set B) only plots values where X is less than a specified cutoff value, and the third curve (data set B) only plots values when X is greater than the cutoff value. The second and third data set are plotted with area fill to shade the tails of the distributions. Remove data set B or C from the graph if you only want to shade one tail. For example, here is the equation used for the first graph (z distribution):

\[ G = \frac{\exp(-0.5 \times X^2)}{\sqrt{2 \times 3.1415926}} \]

\[ A \times Y = G \]

\[ B \times Y = IF(X<-z, G, 0) \]

\[ C \times Y = IF(X>z, G, 0) \]

Change the numbers of degrees of freedom and the cutoff values (for shading) in the Info sheet. This demonstrates how values entered into an info sheet can be 'hooked' to constants used in analyses.

### 3.8.3 Plotting a binomial or Poisson distribution

**How to plot a binomial or Poisson distribution**

![Histograms of binomial and Poisson distributions](image)

**Poisson. True average number (lamda) = 5.3**

**Binomial probability = 0.5**

**n = 12**

**Number of events or objects**

**Number of successes**

[Download the Prism file.](#)

To modify this file, change the value of lamda (for Poission) or the probability, n, and cutoff (Binomial) in the Info sheet. Enter new values there, and the graph updates. This is a good example of the usefulness of hooking an info constant to an analysis.

If you want to recreate graphs like these, keep in mind these points:
• As its name suggests, the analysis 'Create a Family of Theoretical Curves' is usually used to create curves, not bar graphs. When you choose the range of X values, specify the appropriate number of 'line segments' (points) so that the X interval equals 1.0. The binomial example on the left created 16 'line segments' starting at X=0 and ending at X=15. The Poisson on the right created 13 'segments' with X starting at 0 and ending at 13.

• The analysis will create a set of line segments (an attempt to create a curve). Click the change type of graph button, or drop the Change menu and choose Graph Type. Then choose the Grouped tab, and then choose interleaved bars.

• The binomial example on the right has two data sets. You don't want them plotted interleaved, as selected in the previous step. Double click to bring up Format Graph, then go to the middle tab, and choose to superimpose the second data set on the first (rather than interleave). And assign it a different color.

Mathematical details

Binomial distribution

The equation for the probability of exactly X successes in N trials, when each trial has probability P of success is:

\[ R = \text{INT}(X+0.5) \]

\[ \text{ExactProb} = (P^R) \times (1-P)^{(N-R)} \] ; exact probability of successes in n trials

\[ \text{NRearrangments} = \exp(\text{gammaln}(N+1) - \text{gammaln}(R+1) - \text{gammaln}(N-R+1)) \]

\[ \text{gamma}(J) = (J-1)! \] factorial, or \((J-1)!\) but factorial is not a function within Prism \n
\[ \text{NRearrangments} = N! / (R! \times (N-R)!) \]

\[ H = \text{ExactProb} \times \text{NRearrangments} \]

\[ Y = IF(X>cutoff, H, 0) \]

Poisson distribution

The **equation for the Poisson distribution** is:

\[ Y = \exp(-1*\text{Lamda}) \times \text{Lamda}^X / \text{gamma}(X+1) \]

Note the **definition of the gamma function**:

\[ \text{gamma}(i) = \text{factorial}(i-1) \]

\[ \text{gamma}(x+1) = \text{factorial}(x) = X! \]
3.9  Fitting a curve without a model

In some circumstances, your goal is simple. You don't care about models, and don't expect best-fit values that you can interpret. Instead, you just want to draw a smooth curve to make a graph look attractive, or to use as a standard curve.

- **Spline and Lowess curves**
- **Using nonlinear regression with an empirical model**

### 3.9.1 Spline and Lowess curves

#### Curve fitting without a model

The term *curve fitting* is more general than *regression*. Your approach to curve fitting depends on your goal.

In some circumstances, you just want to draw a smooth curve to make a graph look attractive, or to use as a standard curve. You don't care about models, and aren't looking for best-fit values that you can interpret.

Prism provides two approaches for fitting a curve without selecting a model. From a table or graph of XY data, click Analyze, and then choose 'Fit spline/LOWESS' from the list of XY analyses.

#### Spline and lowess curves

A **lowess** curve follows the trend of the data and tends to be a bit jagged. Lowess curves can be helpful when the data progresses monotonically, but are less helpful when there are peaks or valleys. Prism lets you choose between coarse, medium and fine lowess curves. The coarse curve (left panel below) shows only the general trend, but obscures the detail. The fine curve (middle panel below) reveals the fine structure of the data, but tends to wiggle a lot. A **cubic spline** curve (right panel below) goes through every data point, bending and twisting as needed.
Prism generates lowess curves using an algorithm adapted from reference 1. Details. Don't select a lowess curve unless you have well over twenty data points. Prism generates the curve as a series of line segments. Enter the number of segments you want, and check the option box if you need to see the XY coordinates of each point, or if you want to use the resulting lowess, point-to-point, or spline curve as a standard curve. Prism always generates a lowess curve with at least four times more line segments than the number of data points, and you cannot reduce the number of segments below this value.

Creating a point-to-point 'curve'

Prism's spline/lowess analysis can also create a point-to-point "curve" -- a series of line segments connecting all your data. Don't create a point-to-point curve just so you can connect points with a line on the graph. You can do that by checking an option on the Format Graph dialog from the Graphs section of your project. Only select the point-to-point analysis if you want to use the point-to-point line as a standard curve, or if you wish to calculate area under that curve.

References


3.9.2 Using nonlinear regression with an empirical model

If your goal is just to plot a smooth curve, without worrying about a model, you have several choices.

Splines created by Prism go through every point, so may wiggle too much.

Lowess curves follow the general trend of the data, but can be too jagged.

An alternative is to use nonlinear regression.
Nonlinear regression requires you pick a model, but you don't have to pay attention to the meaning of the model or the value of the parameters. Instead, you can pick a model empirically and judge it solely on the appearance of the curve. In this case, you are using nonlinear regression as a tool to create a smooth curve, and not as a method to analyze data.

If you use nonlinear regression in this way, you can experiment with any model you want. But first try fitting polynomial models, which are very general (and never give fitting problems due to poor initial values). If the curve strays too far from the trend of the data, pick a higher order model. If the polynomial curve wiggles too much, pick a lower order model.
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