

Package: pbm (via r-universe)

September 11, 2024

Title Protein Binding Models

Version 1.2.1

Description Binding models which are useful when analysing protein-ligand interactions by techniques such as Biolayer Interferometry (BLI) or Surface Plasmon Resonance (SPR). Naman B. Shah, Thomas M. Duncan (2014) <[doi:10.3791/51383](https://doi.org/10.3791/51383)>. Hoang H. Nguyen et al. (2015) <[doi:10.3390/s150510481](https://doi.org/10.3390/s150510481)>. After initial binding parameters are known, binding curves can be simulated and parameters can be varied. The models within this package may also be used to fit a curve to measured binding data using non-linear regression.

Depends R (>= 3.4.4)

License MIT + file LICENSE

Encoding UTF-8

URL <https://github.com/jonathanrd/pbm>

BugReports <https://github.com/jonathanrd/pbm/issues>

RoxygenNote 7.1.1

Suggests testthat, knitr, rmarkdown, ggplot2, gridExtra

VignetteBuilder knitr

Repository <https://jonathanrd.r-universe.dev>

RemoteUrl <https://github.com/jonathanrd/pbm>

RemoteRef HEAD

RemoteSha f227039c76f0471cbbfe689dc38bbbbd4586a0c92

Contents

| | |
|-----------------------|---|
| binding1to1 | 2 |
| binding2to1 | 2 |
| req | 4 |
| tteq | 4 |

Index

5

binding1to1*Generate a 1:1 Binding Curve***Description**

Returns a response value for given parameters at time, t.

Usage

```
binding1to1(t, t0, conc, kon, koff, rmax, drift = 0, offset = 0, doffset = 0)
```

Arguments

| | |
|---------|---|
| t | Time. |
| t0 | Time of dissociation. |
| conc | Analyte concentration. |
| kon | Kon binding constant. |
| koff | Koff binding constant. |
| rmax | Maximum response, Rmax. |
| drift | Optional. Parameter to add a linear baseline drift. |
| offset | Optional. Applies a global offset to the response value. |
| doffset | Optional. Applies an offset at the start of dissociation. |

Examples

```
time <- seq(1,2000)
curve <- binding1to1(time,1000,6e-9,1000,0.01,0.6)
plot(curve)
```

binding2to1*Generate a 2:1 Binding Curve***Description**

Returns a response value for given parameters at time, t.

Usage

```
binding2to1(  
  t,  
  t0,  
  conc,  
  kon1,  
  koff1,  
  rmax1,  
  kon2,  
  koff2,  
  rmax2,  
  drift = 0,  
  offset = 0,  
  doffset = 0  
)
```

Arguments

| | |
|---------|---|
| t | Time. |
| t0 | Time of dissociation. |
| conc | Analyte concentration. |
| kon1 | Kon binding constant for first component. |
| koff1 | Koff binding constant for first component. |
| rmax1 | Maximum response, Rmax, for first component. |
| kon2 | Kon binding constant for second component. |
| koff2 | Koff binding constant for second component. |
| rmax2 | Maximum response, Rmax, for second component. |
| drift | Optional. Parameter to add a linear baseline drift. |
| offset | Optional. Applies a global offset to the response value. |
| doffset | Optional. Applies an offset at the start of dissociation. |

Examples

```
time <- seq(1,2000)  
curve <- binding2to1(time,1000,900e-9,10000,0.01,0.4,2000,0.0003,0.5)  
plot(curve)
```

| | |
|------------------|--------------------------------|
| <code>req</code> | <i>Response at equilibrium</i> |
|------------------|--------------------------------|

Description

Returns the response value at equilibrium from concentration, Rmax and KD.

Usage

```
req(conc, rmax, kd)
```

Arguments

| | |
|-------------------|------------------------------------|
| <code>conc</code> | Analyte concentration. |
| <code>rmax</code> | Maximum response. |
| <code>kd</code> | Equilibrium dissociation constant. |

Examples

```
req(6e-7, 1.2, 6e-7)
```

| | |
|-------------------|----------------------------|
| <code>tteq</code> | <i>Time to Equilibrium</i> |
|-------------------|----------------------------|

Description

Returns the time taken to reach 95% equilibrium.

Usage

```
tteq(conc, kon, koff, theta = 0.95)
```

Arguments

| | |
|--------------------|------------------------|
| <code>conc</code> | Analyte concentration. |
| <code>kon</code> | Kon binding constant. |
| <code>koff</code> | Koff binding constant. |
| <code>theta</code> | Default 0.95. |

Examples

```
tteq(6e-7, 20000, 0.01)
```

Index

- * **binding1to1**
 - binding1to1, [2](#)
- * **binding2to1**
 - binding2to1, [2](#)

binding1to1, [2](#)
binding2to1, [2](#)

req, [4](#)

tteq, [4](#)