

AQT0794 - BMX Assay Validation

PhosphoSens®-Kinetic Assay Format

Outline for this Study



PhosphoSens-Kinetic Assay Validation

Enzyme Source, Construct, and Lot Information:

Carna BMX (08-179/13CBS-0583K) amino acids full length; N-terminal GST tag

Reference Compound Information:

Staurosporine MedChemExpress (Cat#/Lot#: HY-15141/125391) CAS No.: 62996-74-1

Experiments to be run:

Enzyme Titration

Sensor Peptide K_m Determination

ATP K_m Determination

DMSO Tolerance Test

Reference Compound IC_{50} Determination at ATP K_m

Enzyme Titration

Reaction Conditions and Set Up

AssayQuant®

Reaction Conditions:

54 mM HEPES, pH 7.5

1mM ATP

1.2 mM DTT

0.012% Brij-35

1% glycerol

0.2 mg/ml BSA

0.55 mM EGTA

10 mM MgCl₂

15 μM AQT0794

0.02, 0.05, 0.08, 0.1, 0.2, 0.4, 0.6, 0.8, 1, 2, 3, and 5 nM BMX

Reaction Set Up:

2 or 2.5 μL 10x Sensor Peptide

14 or 17.5 μL Reaction Mix with ATP & DTT

4 or 5 μL 1x EDB or Kinase dilutions (5x in EDB)

20 or 25 μL Final reaction volume

Reactions were run at 30°C for 240 minutes in either Corning, low volume 384-well, white flat round bottom polystyrene NBS microplates (Cat. #3824) at 20 or 25 μ L final well volume or in in PerkinElmer, ProxiPlate-384 Plus, white shallow well microplates (Cat. #6008280) at 20 μ L final well volume after sealing using optically-clear adhesive film (TopSealA-Plus plate seal, PerkinElmer [Cat. #6050185]) in a Biotek Synergy Neo 2 microplate reader with excitation (360 nm) and emission (485 nm) wavelengths.

Notes:

Enzyme Titration



Progress Curves

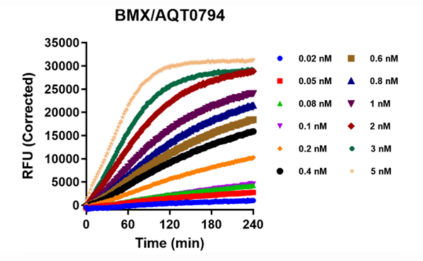
Complete Progress Curves

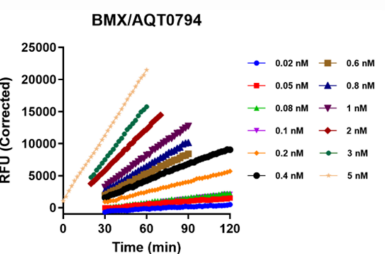
Linear

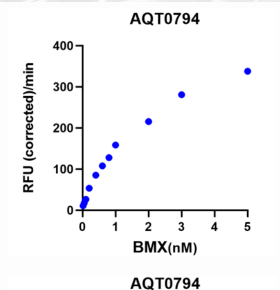
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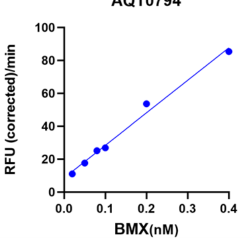
Progress

Curves









Linear Range

Enzyme Titration



Reaction Rate Table

| France Cons (nD4) | Normalized | Normalized Rate |
|-------------------|-------------------------------|----------------------------|
| Enzyme Conc. (nM) | Reaction Rate (RFU/pmole/min) | Stnd Error (RFU/pmole/min) |
| 0.02 | 27,725 | 781 |
| 0.05 | 17,670 | 366 |
| 0.08 | 15,750 | 164 |
| 0.1 | 13,495 | 161 |
| 0.2 | 13,418 | 81 |
| 0.4 | 10,678 | 69 |
| 0.6 | 9,008 | 58 |
| 0.8 | 8,000 | 41 |
| 1 | 7,935 | 49 |
| 2 | 5,395 | 20 |
| 3 | 4,683 | 26 |
| 5 | 3,380 | 14 |

The reaction is linear from 0.02 - 0.4 nM

Sensor Peptide K_m Determination



Reaction Conditions and Set Up

Reaction Conditions:

54 mM HEPES, pH 7.5

1mM ATP

1.2 mM DTT

0.012% Brij-35

1%glycerol

0.2 mg/ml BSA

0.55 mM EGTA

10 mM MgCl₂

1, 2, 3, 5, 7, 10, 15, 20, 30, 50, 70, and 100 μM AQT0794

3 nM BMX

Reaction Set Up:

2 or 2.5 μL 10x Sensor Peptide

14 or 17.5 μL Reaction Mix with ATP & DTT

4 or 5 μL 1x EDB or Kinase dilutions (5x in EDB)

20 or 25 μL Final reaction volume

Reactions were run at 30°C for 240 minutes in either Corning, low volume 384-well, white flat round bottom polystyrene NBS microplates (Cat. #3824) at 25 μ L final well volume or in in PerkinElmer, ProxiPlate-384 Plus, white shallow well microplates (Cat. #6008280) at 20 μ L final well volume after sealing using optically-clear adhesive film (TopSealA-Plus plate seal, PerkinElmer [Cat. #6050185]) in a Biotek Synergy Neo 2 microplate reader with excitation (360 nm) and emission (485 nm) wavelengths.

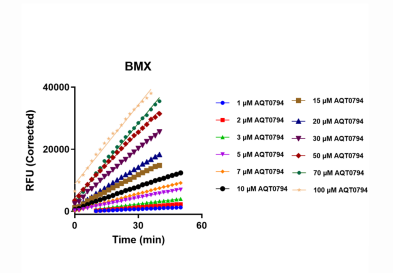
Notes:

Sensor Peptide K_m Determination

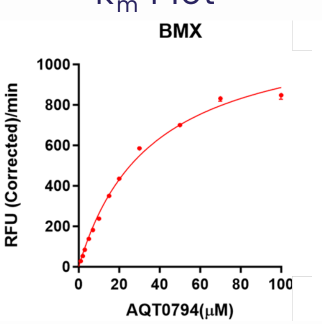


Titration Curves and K_m Plot and Table

Sensor Peptide Titration Curves



Sensor Peptide K_m Plot



Sensor Peptide K_m is 36 µM

Sensor Peptide K_m Table

| Michaelis-Menten | |
|-----------------------------|----------------|
| Best-fit values | |
| Vmax | 1203 |
| Km | 35.83 |
| Std. Error | |
| Vmax | 46.23 |
| Km | 3.061 |
| 95% CI (profile likelihood) | |
| Vmax | 1110 to 1312 |
| Km | 29.89 to 43.21 |
| Goodness of Fit | |
| Degrees of Freedom | 10 |
| R squared | 0.9948 |
| | |

ATP K_m Determination



Reaction Conditions and Set Up

Reaction Conditions:

54 mM HEPES, pH 7.5

0, 2.0, 3.9, 7.8, 16, 31, 63, 125, 250, 500, 1000, and 2000 μM ATP

1.2 mM DTT

0.012% Brij-35

1%glycerol

0.2 mg/ml BSA

0.55 mM EGTA

10 mM MgCl₂

15 μM AQT0794

3 nM BMX

Reaction Set Up:

2 or 2.5 μL 10x Sensor Peptide

14 or 17.5 μL Reaction Mix with ATP & DTT

4 or 5 μL 1x EDB or Kinase dilutions (5x in EDB)

20 or 25 μL Final reaction volume

Reactions were run at 30°C for 240 minutes in either Corning, low volume 384-well, white flat round bottom polystyrene NBS microplates (Cat. #3824) at 25 μ L final well volume or in in PerkinElmer, ProxiPlate-384 Plus, white shallow well microplates (Cat. #6008280) at 20 μ L final well volume after sealing using optically-clear adhesive film (TopSealA-Plus plate seal, PerkinElmer [Cat. #6050185]) in a Biotek Synergy Neo 2 microplate reader with excitation (360 nm) and emission (485 nm) wavelengths.

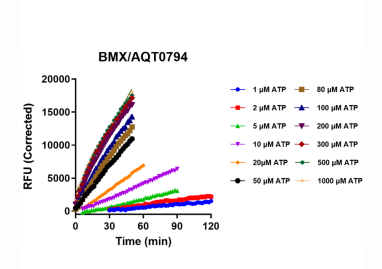
Notes:

ATP K_m Determination

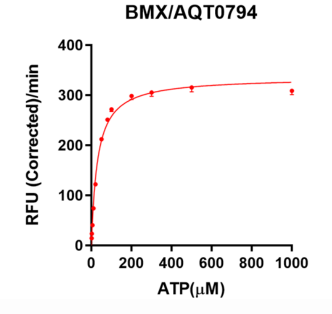


Titration Curves and K_m Plot and Table

ATP Titration Curves



ATP K_m Plot



ATP K_m is 30 μM

ATP K_m Table

| Michaelis-Menten | |
|-----------------------------|----------------|
| Best-fit values | |
| Vmax | 335.6 |
| Km | 30.24 |
| Std. Error | |
| Vmax | 6.090 |
| Km | 2.437 |
| 95% CI (profile likelihood) | |
| Vmax | 322.6 to 349.1 |
| Km | 25.45 to 35.84 |
| Goodness of Fit | |
| Degrees of Freedom | 10 |
| R squared | 0.9945 |
| · | • |

DMSO Tolerance Test



Reaction Conditions and Set Up

Reaction Conditions:

54 mM HEPES, pH 7.5

1 mM ATP

1.2 mM DTT

0.012% Brij-35

1% glycerol

0.2 mg/ml BSA

0.55 mM EGTA

10 mM MgCl₂

0, .01, .02, .04, .08, .16, .31, .63, 1.3, 2.5, 5.0, and 10% DMSO

15 μM AQT0794

3 nM BMX

Reaction Set Up:

2 or 2.5 μL 10x DMSO dilutions

14 or 17.5 μL Reaction Mix with Sensor Peptide, ATP & DTT

4 or 5 μL 1x EDB or Kinase dilutions (5x in EDB)

20 or 25 μL Final reaction volume

Reactions were run at 30°C for 240 minutes in either Corning, low volume 384-well, white flat round bottom polystyrene NBS microplates (Cat. #3824) at 25 μ L final well volume or in in PerkinElmer, ProxiPlate-384 Plus, white shallow well microplates (Cat. #6008280) at 20 μ L final well volume after sealing using optically-clear adhesive film (TopSealA-Plus plate seal, PerkinElmer [Cat. #6050185]) in a Biotek Synergy Neo 2 microplate reader with excitation (360 nm) and emission (485 nm) wavelengths.

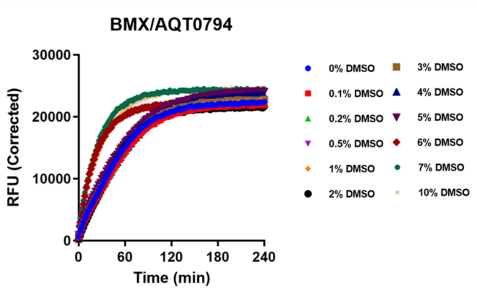
Notes:

DMSO Tolerance Test

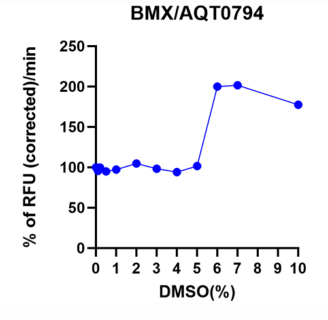


Titration Curves and Inhibition Plot





Reaction Rate vs [DMSO] Plot



No change in enzyme activity out to 5% DMSO

IC₅₀ Determination

AssayQuant®

Reaction Conditions and Set Up

Reaction Conditions:

54 mM HEPES, pH 7.5

ATP at K_m

1.2 mM DTT

0.012% Brij-35

1%glycerol

0.2 mg/ml BSA

0.55 mM EGTA

10 mM MgCl₂

1% DMSO

15 μM AQT0794

2 nM BMX

0.1 mM Staurosporine was serially diluted (3-fold, 11-point) in 100%DMSO. The series was then diluted 10-fold into BSA (with a final concentration of 0.2 mg/mL BSA in 10% DMSO) to prepare the 10x compound stocks.

Reaction Set Up:

16 μL Reaction Mix with Sensor Peptide and Inhibitor

 $4 \mu L$ 1x EDB or Kinase dilutions (5x in EDB)

20 μL Final reaction volume

Reactions were run at 30°C for 240 minutes in either Corning, low volume 384-well, white flat round bottom polystyrene NBS microplates (Cat. #3824) at 20 μ L final well volume or in in PerkinElmer, ProxiPlate-384 Plus, white shallow well microplates (Cat. #6008280) at 20 μ L final well volume after sealing using optically-clear adhesive film (TopSealA-Plus plate seal, PerkinElmer [Cat. #6050185]) in a Biotek Synergy Neo 2 microplate reader with excitation (360 nm) and emission (485 nm) wavelengths.

Inhibitors are added via direct (0.4 μ L of 50X stock in 100% DMSO) or intermediate dilutions (2.0 μ L of 10X stock in 10% DMSO).

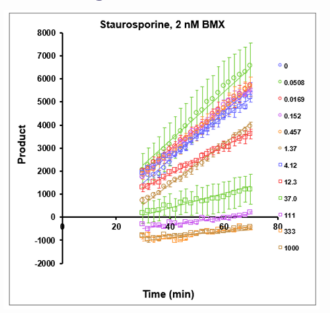
Notes:

IC₅₀ Determination

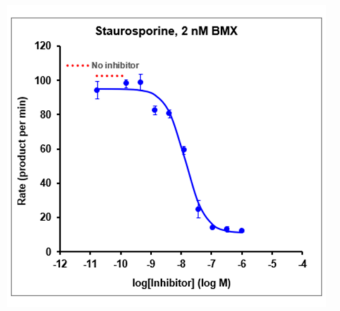


Progress Curves and IC₅₀ Curves and Table

Linear Region of Progress Curves



IC₅₀ Curve



IC₅₀ Table

| Parameter | Value |
|---------------------|--------|
| Bottom | 10.7 |
| Тор | 94.9 |
| log IC50 | -7.87 |
| IC50 (nM) | 13.54 |
| Ki (nM) | 6.77 |
| Slope | -1.347 |
| R squared | 0.988 |
| C50 approx SE (nM) | 0.61 |
| 50% inhibition (nM) | 16.37 |

The Y-axis label is RFU/min.

Staurosporine IC₅₀ at ATP K_m is 14 nM

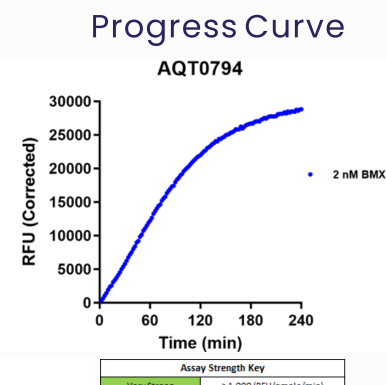
Summary



Assay Validation Results and Progress Curve and Assay Strength at 1 mM ATP

| Experiment | Result |
|--|---------------|
| Enzyme Titration Linear Range | 0.02 - 0.4 nM |
| Sensor Peptide K _m Value | 36 μΜ |
| ATP K _m Value | 30 μΜ |
| DMSO Tolerance (highest % recommended) | 5 |
| Staurosporine IC ₅₀ Determination at ATP K _m | 14 nM |

| Kinase Name | Conc. (nM) | Sox-based Substrate Name | Normalized Reaction Rate (RFU/pmole/min) | Normalized Rate StndError (RFU/pmole/min) |
|-------------|------------|-----------------------------|--|---|
| вмх | 2 | AQT0794 | 5,395 | 20 |



| Assay Strength Key | |
|--------------------|----------------------------|
| Very Strong | >1,000 (RFU/pmole/min) |
| Strong | 300 to 999 (RFU/pmole/min) |
| Moderate | 100 to 299 (RFU/pmole/min) |
| Weak | 30 to 99 (RFU/pmole/min) |

Under the conditions utilized for this experiment, the assay is Very Strong