

# AQT1030 - VRK1 Assay Validation

PhosphoSens®-Kinetic Assay Format

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# **Outline for this Study**



PhosphoSens-Kinetic Assay Validation

### **Enzyme Source, Construct, and Lot Information:**

Active Motif VRK1 (Cat#/Lot#: 81096/32317001) amino acids 1-396(end), N-terminal FLAG tag

#### **Reference Compound Information:**

Staurosporine MedChemExpress (Cat#/Lot#: HY-15141/125391)

#### **Experiments to be run:**

**Enzyme Titration** 

Sensor Peptide K<sub>m</sub> Determination

ATP K<sub>m</sub> Determination

**DMSO Tolerance Test** 

Reference Compound IC<sub>50</sub> Determination at ATPK<sub>m</sub>

# **Enzyme Titration**

### **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 MgCl<sub>2</sub> 15 μM AQT1030

0, 0.08, 0.16, 0.31, 0.63, 1.3, 2.5, 5, 10, 20, 40, and 80 nM VRK1

# Reaction Set Up:2 or 2.5 μL10x Sensor Peptide14 or 17.5 μLReaction Mix with A4 or 5 μL1x EDB or Kinase dil

Reaction Mix with ATP & DTT 1x EDB or Kinase dilutions (5x in EDB) 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  $\mu$ L final well volume or in in PerkinElmer, ProxiPlate-384 Plus, white shallow well microplates (Cat. #6008280) at 20  $\mu$ 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:

20 or 25 µL





# **Enzyme Titration**

### **Progress Curves**



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# **Enzyme Titration**



### **Reaction Rate Table**

Enzyme Conc. (nM)	Reaction Rate (RFU/min)	Normalized Reaction Rate (RFU/pmole/min)
0.08	-0.42	-261
0.16	-0.60	-186
0.33	0.91	138
0.65	3.3	252
1.3	7.7	295
2.5	14	284
5	25	246
10	48	239
20	80	200
40	134	168
80	216	135

### The reaction is linear from 0.65 - 10 nM

# Sensor Peptide K<sub>m</sub> 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 MgCl<sub>2</sub> 0, .6, 1.2, 2.3, 4.7, 9.4, 18.8, 37.5, 75, 150, 300, and 600 μM AQT1030 20 nM VRK1

# Reaction Set Up:2 or 2.5 μL10x Sensor Peptide14 or 17.5 μLReaction Mix with ATP & DTT4 or 5 μL1x EDB or Kinase dilutions (5x in EDB)20 or 25 μLFinal 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  $\mu$ L final well volume or in in PerkinElmer, ProxiPlate-384 Plus, white shallow well microplates (Cat. #6008280) at 20  $\mu$ 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**<sub>m</sub> Determination

Titration Curves and K<sub>m</sub> Plot and Table

Sensor Peptide Titration Curves





Sensor Peptide

K<sub>m</sub> Plot

## Sensor Peptide K<sub>m</sub> Table

Parameter	Value	Approx SE
Vmax (RFU per min)	315.7	6.2
Vmax (RFU per pmol per min)	789	16
Km (μM)	187.7	16.6
R squared	0.980	

### Sensor Peptide K<sub>m</sub> is 188 µM



# ATP K<sub>m</sub> 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 $\mu M$ ATP
1.2 mM DTT
0.012% Brij-35
1%glycerol
0.2 mg/ml BSA
0.55 mM EGTA
10 mM MgCl <sub>2</sub>
15 μM AQT1030
20 nM VRK1

# Reaction Set Up:<br/>2 or 2.5 μL10x Sensor Peptide14 or 17.5 μLReaction Mix with ATP & DTT4 or 5 μL1x EDB or Kinase dilutions (5x in EDB)20 or 25 μLFinal 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  $\mu$ L final well volume or in in PerkinElmer, ProxiPlate-384 Plus, white shallow well microplates (Cat. #6008280) at 20  $\mu$ 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<sub>m</sub> Determination**

Titration Curves and K<sub>m</sub> Plot and Table

**2000** 

01000

○ 500

250

0130

**O63** 

□31 □16

3.9

2.0

50 <sub>07.8</sub>

ATP Titration Curves

20 nM VRK1, 15 µM AQT1030

1600

1400

1200

1000

800

600

400

200

-200

-400

-600

Product (RFU)



ATP K<sub>m</sub> Plot

## ATP K<sub>m</sub> Table

Parameter	Value	Approx SE
Vmax (RFU per min)	30.7	1.1
Vmax (RFU per pmol per min)	77	3
Km (μM)	51.3	6.5
R squared	0.995	

### ATP K<sub>m</sub> is 51 µM

Time (min)





# **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 MgCl<sub>2</sub> 0, .01, .02, .04, .08, .16, .31, .63, 1.3, 2.5, 5.0, and 10% DMSO 15 μM AQT1030 20 nM VRK1 Reaction Set Up:2 or 2.5 μL10x DMSO dilutions14 or 17.5 μLReaction Mix with Sensor Peptide, ATP & DTT4 or 5 μL1x EDB or Kinase dilutions (5x in EDB)20 or 25 μLFinal 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  $\mu$ L final well volume or in in PerkinElmer, ProxiPlate-384 Plus, white shallow well microplates (Cat. #6008280) at 20  $\mu$ 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** 

## Complete Progress Curves







No change in enzyme activity out to 3% DMSO, but significant activation (150%) at 5% and 10% DMSO. This has been seen with other kinases.

# IC<sub>50</sub> Determination

### **Reaction Conditions and Set Up**

#### **Reaction Conditions:**

54 mM HEPES, pH 7.5 ATP at K<sub>m</sub> 1.2 mM DTT 0.012% Brij-35 1%glycerol 0.2 mg/ml BSA 0.55 mM EGTA 10 mM MgCl<sub>2</sub> 2% DMSO

 $15 \,\mu\text{M}$  AQT1030

20 nM VRK1

0, 0.169, 0.508, 1.52, 4.57, 13.7, 41.2, 123, 370, 1110, 3330, 10000 nM Staurosporine

## Reaction Set Up:

16 μL Reaction Mix with Sensor Peptide and Inhibitor
<u>4 μL</u>
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  $\mu$ L final well volume or in in PerkinElmer, ProxiPlate-384 Plus, white shallow well microplates (Cat. #6008280) at 20  $\mu$ 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  $\mu$ L of 50X stock in 100% DMSO) or intermediate dilutions (2.0  $\mu$ L of 10X stock in 10% DMSO).

#### Notes:



# IC<sub>50</sub> Determination

00

0.508

00.169

01.52

4 57

0137

**141 2** 

**123** 

370

1110

3330

150 10000



Progress Curves and IC<sub>50</sub> Curves and Table

Inhibitor Titration Progress Curves

Staurosporine, 20 nM Active Motif VRK1

50

Time (min)

100

3000

2500

2000

1500

1000

500

-500

Product

## Linear Region of Progress Curves

Staurosporine, 20 nM Active Motif VRK1 2000 00 1500 0.508 00.169 01.52 1000 Product 04.57 013.7 **41.2** 500 **123** 370 1110 *۸* 60 80<sub>3330</sub> 10000 -500 Time (min)

IC<sub>50</sub> Curve

Staurosporine, 20 nM Active Motif VRK1

log[Inhibitor] (log M)

30

25

20

15

10

5

0 + -11 ••••• No inhibitor

-10

per min)

Rate (product

IC<sub>50</sub> Table

Parameter	Value
Bottom	0.0
Тор	20.8
log IC50	-5.11
IC50 (nM)	7805.06
Ki (nM)	3902.53
Slope	-0.954
<b>R</b> squared	0.970
C50 approx SE (nM)	Large
50% inhibition (nM)	7805.06



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Staurosporine IC<sub>50</sub> at ATP K<sub>m</sub> is 7800 nM

# Summary



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

Experiment	Result	Progress Curve
Enzyme Titration Linear Range	10 - 40 nM	AQT1030
Sensor Peptide K <sub>m</sub> Value	188 µM	10000 T T T T T T T T T T T T T T T T T
ATP K <sub>m</sub> Value	51 μΜ	6000 - 50 9 4000 - 20 nM VRK1(Active Motif)
DMSOTolerance (highest % recommended)	10%	
Staurosporine $\rm IC_{50}$ Determination at ATP $\rm K_m$	7800 nM	を 0 60 120 180 240 Time [min]

	Conc. (nM) Sox-base Substrate N	Sox-based Substrate Name (RFU/pmole/min)	Normalized	Normalized Pate	
Kinasa Nama			Reaction Rate	Stad Favor	
Kinase Name			Substrate Name	(RFU/pmole/min)	
			(RFU/pmole/min)		(RFU/pmole/min)
VRK1	20	AQT1030	200	2.6	

### Under the conditions utilized for this experiment, the assay is Moderate