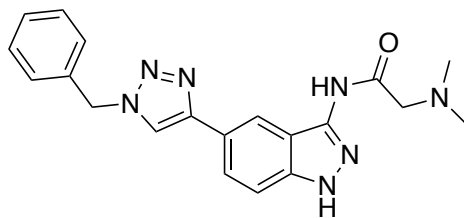


# CSNK1G2



AKI00000062a

**Chemical Name:**

*N*-(5-(1-benzyl-1*H*-1,2,3-triazol-4-yl)-1*H*-indazol-3-yl)-2-(dimethylamino)acetamide

**CHEBI:** 143112

**Smile String:**

O=C(CN(C)C)NC1=NNC2=CC=C(C3=CN(CC4=CC=CC=C4)N=N3)C=C21

**Chemical Formula:** C<sub>20</sub>H<sub>21</sub>N<sub>7</sub>O

**Molecular Weight:** 375.44

**cLogP:** 1.8746

**Source:** SGC-UNC

**Reference:**

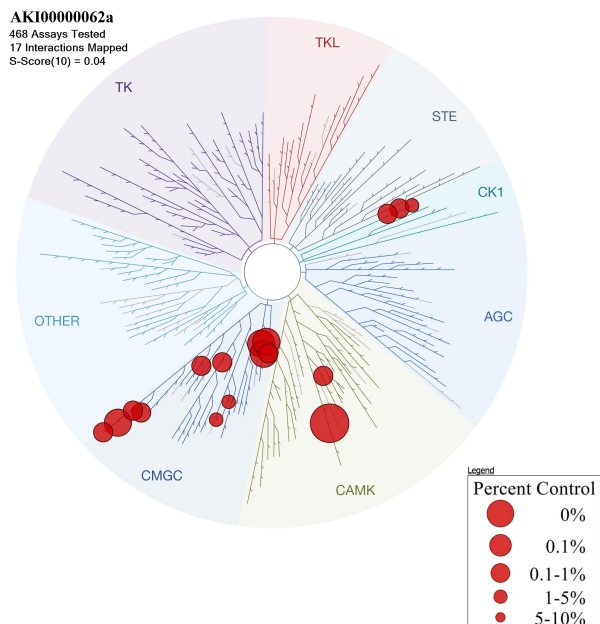
US 20090203690

## Biochemical profiling

DiscoverX (403 wild-type human kinases)

**S<sub>10</sub> (1 μM): 0.042**

(17 kinases < 10% control)



Kinase	% Control @ 1uM
CAMK1B	0
CDKL5	0.2
PCTK1	0.2
GSK3B	0.4
GSK3A	0.9
CSNK1G2	1
CDK7	1.2
PCTK3	1.2
PCTK2	1.4
ICK	1.6
DCAMKL3	1.9
ERK8	2.5
CSNK1G3	3.3
PFTK1	3.5
CSNK1G1	5.2
DYRK1B	6
DYRK2	7.2

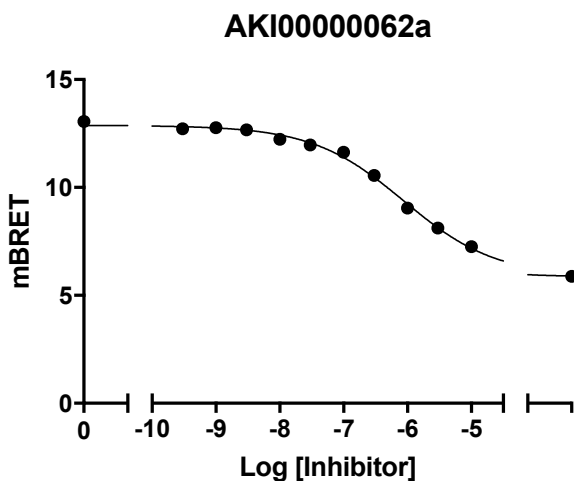
a. Treemap of DiscoverX KINOMEScan data.

b. List of kinases inhibited < 10% control

## Cellular target engagement in HEK293 cells

NLuc-CSNK1G2 (N term)

CSNK1G2 IC<sub>50</sub> = 876 nM



Cellular target engagement of AKI00000062a with CSNK1G2