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Inhibitor catalog







Inhibitors

Selleck Chemicals supplies over 3,000 inhibitors used in the study of cell signaling pathways.



Product Citations

Selleck products have been cited in more than **27000** studies from various **SCI** journals. (**Cell, Nature, Science: 77** studies)

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Compound Libraries



Customize your library by selecting compounds of interest.

Selleck is a Licensed Supplier of Pfizer Compounds



In 2013, Selleck became a licensed supplier of Pfizer pharmaceuticals. This has granted our customers access to Pfizer's exclusive and high quality compounds. Purchased individually or as a library, these compounds have a wide range of applications in preclinical research of human diseases.

• All bioactive compounds are licensed by Pfizer and have been marketed and/or have been clinically demonstrated to be safe and efficacious in humans.

• Compounds span a range of potential uses: from anti-cancer compounds (e.g. Bosutinib) to a glycylcycline antiobiotic (e.g. Tigecycline) to combat the growing prevalence of antibiotic resistance.

• Reliability Guarantee: all Pfizer licensedcompounds are developed, and validated by Pfizer, and some even manufactured by Pfizer Quality Assurance: all compounds are validated using NMR and HPLC.

Detailed preclinical research data and safety information available.

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РІЗК
mTOR
Akt
GSK-3
ATM/ATR
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AMPK
DNA-PK
MELK

Epigenetics

HDAC
PARP
JAK
Pim
HIF
Aurora Kinase 27
Sirtuin
Epigenetic Reader Domain
Histone Acetyltransferase
DNA Methyltransferase
Histone Methyltransferase
Histone Demethylase

Protein Tyrosine Kinase

VEGFR
EGFR
PDGFR
c-Met
HER2
IGF-1R
FLT3
FGFR
c-Kit
ALK
Trk Receptor
Ephrin Receptor 48
CSF-1R
TAM Receptor

Angiogenesis

VEGFR
JAK
EGFR
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GluR
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AChR

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wicrobiology	HCV Protease 128
microbiology-	HCV Protease 128 HIV Protease 128
microbiology-	HCV Protease
microbiology-	HCV Protease128HIV Protease128Integrase128Reverse Transcriptase129
microbiology-	HCV Protease128HIV Protease128Integrase128Reverse Transcriptase129CCR130
Microbiology-	HCV Protease128HIV Protease128Integrase128Reverse Transcriptase129CCR130Antifection130

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FXR
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Phosphatase 132
NADPH Oxidase 132
PTEN
Others

PD-1/PD-L1 Inhibitory Antibodies

A2002 Nivolumab

Nivolumab is a genetically engineered, fully human immunoglobulin (Ig) G4 monoclonal antibody directed against the negative immunoregulatory human cell surface receptor programmed death-1 (PD-1,PCD-1) with immune checkpoint inhibitory and antineoplastic activities. Size 5 mg

A2004 Atezolizumab Atezolizumab is a fully humanized, IgG1 monoclonal antibody that blocks the interaction of PD-L1 with both PD-1 and B7.1, but not the interaction of PD-L2 with PD-1

Size 5 mg

A2005 Pembrolizumab

Pembrolizumab is a potent, highly selective, fully humanized immunoglobulin (Ig) G4-kappa monoclonal antibody against PD-1 with potential immune checkpoint inhibitory and antineoplastic activities. Size 5 ma

A2013 Durvalumab

Durvalumab is a human immunoglobulin G1 kappa (IgG1k) monoclonal antibody that blocks the interaction of programmed cell death ligand 1 (PD-L1) with the PD-1 and CD80 (B7.1) molecules.

Size 5 mg

A2014 BMS-936559

BMS-936559 is a fully human IgG4 mAb to PD-L1 and inhibits the binding of PD-L1 to both PD-1 and CD80 (a ligand of CTLA-4). Size 5 mg

A2015 Avelumab

Avelumab is a human anti-PD-L1 IgG1 monoclonal antibody with antineoplastic actions

Inhibitory Antibodies Size 5 mg

A2016 SHR-1210

SHR-1210 is a humanized anti-PD-1 IgG4 antibody that blocks the binding of PD-L1 and PD-L2 to PD-1 with immune checkpoint inhibitory and antineoplastic activities.

Size 5 mg

A2017 PDR001

PDR001 is a humanized anti-PD-1 IgG4 antibody that blocks the binding of PD-L1 and PD-L2 to PD-1. It binds to PD-1 with high affinity and inhibits the biological activity of PD-1.

Size 5 mg

A2018 Lambrolizumab

Lambrolizumab is a humanized monoclonal IgG4 antibody against PD-1 that blocks the interaction between PD-1 and its ligands, PD-L1 and PD-L2.

Size 5 mg

Other Inhibitory Antibodies

A2000 Cetuximab

Cetuximab, a novel molecular-targeted agent, is an inhibitor of EGFR monoclonal humanized antibody interacting with the extracellular binding site of EGFR to block ligand stimulation.

Size 5 mg

A2006 Bevacizumab

Bevacizumab is a humanized anti-VEGF monoclonal antibody which binds to and neutralizes all human VEGF-A isoforms and bioactive proteolytic fragments

Size 5 mg

A2007 Trastuzumab

Trastuzumab is a humanized, recombinant monoclonal antibody that binds to the extracellular domain of HER2. Size 5 mg

A2008 Pertuzumah

Pertuzumab, a humanized monoclonal antibody and the first in the class of agents called the HER2 dimerization inhibitors, impairs the ability of HER2 to bind to other members of the HER family. Size 5 mg

A2011 Sarilumab

Sarilumab is a fully human anti-IL-6Ra mAb that binds membrane-bound and soluble human IL-6Rg with high affinity Size 5 mg

A2012 Tocilizumab

Tocilizumab is a humanized monoclonal antibody that binds to the interleukin-6 receptor Size 5 mg

A2019 Abatacept

Abatacept, a selective T-cell costimulation modulator, is a soluble fusion protein comprising the extracellular domain of human cytotoxic T-lymphocyte-associated antigen-4 (CTLA4) linked to the modified Fc (hinge, CH2 and CH3 domains) portion of human immunoglobulin G1 (CTLA-Ig).

A2020 Ustekinumab

Size 5 mg

Ustekinumab is an anti IL-12/23 IgG1 kappa human monoclonal antibody that targets the p40 subunit shared by two cytokines, interleukin (IL)-12 and 23, prevents their interaction with the receptor, thereby blocking subsequent signaling, differentiation and cytokine production central to inflammatory diseases. Size 5 mg

A2021 Obinutuzumab

5 mg

Obinutuzumab (GA101) is a novel, type II, glycoengineered, humanized anti-CD20 monoclonal antibody. Size

A2022 Vedolizumab

Vedolizumab is a humanized monoclonal antibody which acts against $\alpha 4\beta 7$ integrin heterodimer and blocks the interaction of $\alpha 4\beta 7$ integrin with MAdCAM-1. It does not inhibit binding at VCAM-1. Size 5 mg

A2023 Eculizumab

Eculizumab is a humanized monoclonal antibody that binds specifically to complement protein C5 with high affinity, preventing its cleavage into C5a and C5b. Size 5 mg

A2024 Daratumumab Daratumumab is a human monoclonal antibody that targets CD38, a cell surface protein that is highly expressed on haematological malignancies Size 5 mg

Bioactive Compound Library Cat.No. L1700

 Over 3300 bioactive chemical compounds for high throughput screening (HTS) and high content screening (HCS)

- Bioactivity and safety confirmed by preclinical research and clinical trials
- Some compounds have been approved by the FDA
- Includes most Selleck inhibitors, APIs, natural products, and chemotherapeutic agents
- Structurally diverse, medicinally active, and cell permeable
- · Rich documentation with structure, IC50, and customer reviews

NMR and HPLC validated to ensure high purity

Size (Pre-dissolve	d in DMSO)	Customize Your Library					
100 µL/well	(10 mM solution)	5	5			II	
2x100 µL/well	(10 mM solution)	Spe Comp	cific Qua	antities	Plate map	Format (Dry/solid or DMSO solution)	



Journals Citing of this Library

Nat Med, 2014, 20(8):954-60

.....

- Oncotarget, 2014, 5(15):6512-25
- Oncotarget, 2015, 6(3):1531-43
- J Biomol Screen, 2015, 20(9):1171-7

cell biology

Compound Library

FDA-approved Drug Library Cat.No. L1300

- A unique collection of 1539 FDA approved drugs for high throughput screening (HTS) and high content screening (HCS)
- · Locate new targets for old drugs
- · Bioactivity and safety confirmed by clinical trials
- All compounds have been approved by FDA
- · Related to oncology, cardiology, anti-inflammatory, immunology, neuropsychiatry, analgesia etc
- Structurally diverse, medicinally active, and cell permeable
- Rich documentation with structure, IC50, and customer reviews
- NMR and HPLC validated to ensure high purity





Journals Citing of this Library

Cancer Res, 2014, 74:1702 Nat Prod Rep, 2014, 31(6):718-29 PLoS One, 2015, 10(6):e0129234 PLoS One, 2015, 10(11):e0143033



Other Compound Libraries

Kinase Inhibitor Library Cat.No. L1200

A unique collection of 504 kinase inhibitors for high throughput screening (HTS) and high content screening (HCS).

Natural Product Library Cat.No. L1400

Over 700 natural products for high throughput screening (HTS) and high content screening (HCS).

Express-Pick Library Cat.No. L3600

A unique collection of 4208 chemical compounds featured different parent nuclei and structural diversities respectively for high throughput screening (HTS) and high content screening (HCS).

Inhibitor Library Cat.No. L1100

A unique collection of 1908 inhibitors for high throughput screening (HTS) and high content screening (HCS).

Epigenetics Compound Library Cat.No. L1900

A unique collection of 210 small molecule modulators with biological activity used for epigenitc research

Target Selective Inhibitor Library Cat.No. L3500

A unique collection of validated bioactive compounds covering over 174 targets.

GPCR Compound Library Cat.No. L2200

A unique collection of 514 GPCR small molecule compound library for GPCR screening.

Anti-cancer Compound Library Cat.No. L3000

A unique collection of 1070 compounds with anti-cancer activity for high throughput screening (HTS) and high content screening (HCS).

Tyrosine Kinase Inhibitor Library Cat.No. L1800

A unique collection of 204 tyrosine kinase inhibitors for high throughput screening (HTS) and high content screening (HCS).

Stem Cell Signaling Compound Library Cat.No. L2100

A unique collection of **101** small molecule inhibitors used for stem cell regulatory and signaling pathway research.

Cambridge Cancer Compound Library Cat.No. L2300

A unique collection of 264 anti-cancer compounds.

Other Compound Libraries

Pfizer Licensed Compound Library Cat.No. L2400

91 bioactive compounds are licensed by Pfizer and have been marketed or clinically proven.

Autophagy Compound Library Cat.No. L2600

A unique collection of 161 autophagy signaling pathway ihibitors.

Ion Channel Ligand Library Cat.No. L2700

A unique collection of **78** ion channel ligands.

Compound Library

PI3K/Akt Inhibitor Library Cat.No. L2800

A unique collection of 127 PI3K signaling pathway inhibitors.

Apoptosis Compound Library Cat.No. L3300

A unique collection of **125** small molecules used for apoptosis research targeting Bcl-2, Caspase, p53, TNF-alpha, Mdm2, survivin, etc.

MAPK Inhibitor Library Cat.No. L3400

A unique collection of 66 small molecule inhibitors used for MAPK signaling research.

Protease Inhibitor Library Cat.No. L2500

A unique collection of **72** small molecule inhibitors used for chemical genomics, high-throughput screening (HTS), and high content screening (HCS).

Anti-infection Compound Library Cat.No. L3100

A unique collection of **338** anti-infective small molecules with biological activity of antibiotics, antifungal drugs, anti-HIV, etc.

Anti-diabetic Compound Library Cat.No. L2900

A unique collection of 33 small molecules affecting the development of diabetes.

Metabolism Compound Library Cat.NO. L3700

A unique collection of 441 small molecule compounds used for metabolic research.

Express-Pick Library (Premium Version) Cat.NO. L5000

A unique collection of **111430** innovative chemical compounds, from the largest pharmaceutical company in the word, features numerous structurally diverse compounds and several alternate compositions, for high throughput screening (HTS) and high content screening (HCS).

PI3K/Akt/mTOR Pathway



PI3K Inhibitors

Inhibitory Selectivity

Inhibitor Name	РІЗК	p110α	p110β	p110ō	p110γ	C2β	Vps34	Other
Dactolisib		++++IC50: 4 nM	++ IC50: 75 nM	+++ IC50: 7 nM	++++IC50: 5 nM			mTOR (p70S6K),ATR
Pictilisib		++++IC50: 3 nM	+++ IC50: 33 nM	++++ IC50: 3 nM	++ IC50: 75 nM			mTOR
LY294002		+ IC ₅₀ : 0.5 μM	+ IC ₅₀ : 0.97 μM	+ IC50: 0.57 μM				
Idelalisib				++++ IC50: 2.5 nM	++ IC50: 89 nM			
Buparlisib		++ IC50: 52 nM	+ IC50: 166 nM	++ IC50: 116 nM	+ IC50: 262 nM		+ IC50: 2.4 μM	mTOR
PI-103		++++IC50: 2 nM	++++ IC50: 3 nM	++++ IC50: 3 nM	+++ IC50: 15 nM			DNA-PK,mTOR
TGX-221			++++ IC50: 5 nM	++ IC50: 0.1 μM				
IC-87114				+ IC ₅₀ : 0.5 μM	+ IC ₅₀ : 29 μM			
Wortmannin	++++ IC50: 3 nM							DNA-PK,ATM,MLCK
XL147 analogue		++ IC50: 39 nM	+ IC50: 383 nM	++ IC50: 36 nM	+++ IC50: 23 nM			
ZSTK474	+++ IC50: 37 nM	+++ IC50: 16 nM	++ IC50: 44 nM	++++ IC50: 4.6 nM	++ IC50: 49 nM			
Alpelisib		++++IC50: 5 nM						
AS-605240		++ IC50: 60 nM	+ IC50: 270 nM	+ IC50: 300 nM	+++ IC50: 8 nM			
PIK-75 HCI		+++ IC50: 5.8 nM		+ IC50: 0.51 μM	++ IC50: 76 nM			DNA-PK
3-Methyladenine					+ IC50: 60 μM		+ IC50: 25 μM	
A66		++ IC50: 32 nM				+ IC50: 462 nM		ΡΙ4Κβ
Voxtalisib Analogue		++ IC50: 39 nM	++ IC50: 113 nM	++ IC50: 43 nM	+++ IC50: 9 nM			DNA-PK,mTOR
PIK-93		++ IC50: 39 nM	+ IC50: 590 nM	++ IC50: 120 nM	+++ IC50: 16 nM	++ IC50: 140 nM	++IC50: 320 nM	PI4KIIIβ,DNA-PK,ATM

PI3K

4

PI3K

Inhibitory Selectivity

	Inhibitor Name	РІЗК	p110α	p110β	p110ō	p110γ	C2β	Vps34	Other
	Omipalisib		++++ Ki: 0.019 nM	++++K: 0.13 nM	++++ Ki: 0.024 nM	++++Ki: 0.06 nM			mTORC1,mTORC2
	PIK-90		+++ ICso: 11 nM	+ IC50: 350 nM	++ IC50: 58 nM	+++ IC50: 18 nM			
5	PF-04691502		++++ Ki: 1.8 nM	++++Ki: 2.1 nM	++++Ki: 1.6 nM	++++Ki: 1.9 nM			P-Akt,P-Akt,mTOR
	AZD6482		+ IC50: 870 nM	+++ IC50: 10 nM	++ IC50: 80 nM				DNA-PK
ş	Apitolisib		++++ IC50: 5 nM	+++ IC50: 27 nM	+++ IC50: 7 nM	+++ IC50: 14 nM			mTOR
2	GSK1059615		++++ IC50: 0.4 nM	++++IC50: 0.6 nM	++++IC50: 2 nM	++++IC50: 5 nM			mTOR
2	Duvelisib			++++K: 1564 pM	++++ Ki: 23 pM	++ Ki: 243 pM			
-	Gedatolisib		++++ IC50: 0.4 nM			+++ IC50: 5.4 nM			mTOR
	TG100-115		+ IC50: 1.3 μM	+ IC ₅₀ : 1.2 μM	+ IC50: 235 nM	++ IC50: 83 nM			
	AS-252424		+ IC50: 935 nM			++ IC50: 33 nM			Casein Kinase 2
	BGT226		++++ IC50: 4 nM	++ IC50: 63 nM		++ IC50: 38 nM			mTOR
	CUDC-907		+++ IC50: 19 nM	++ IC50: 54 nM	+++ IC50: 39 nM				HDAC1,HDAC3,HDAC10
	PIK-294			+ IC50: 490 nM	+++ IC50: 10 nM	+ IC50: 160 nM			
	AS-604850		+ IC50: 4.5 μM			+ ICso: 0.25 μM			
	Copanlisib		++++ IC50: 0.5 nM	++++IC50: 3.7 nM	++++IC50: 0.7 nM	++++IC50: 6.4 nM			
	YM201636		+ ICso: 3.3 μM						PIKfyve
	CH5132799		+++ ICso: 14 nM	++ IC50: 0.12 μM	+ IC50: 0.50 μM	++ IC50: 36 nM			
	PIK-293				+ IC ₅₀ : 0.24 μM	+ IC ₅₀ : 10 μM			
	PKI-402		++++ IC50: 2 nM	+++ IC50: 7 nM	+++ IC50; 14 nM	+++ IC∞: 16 nM			mTOR
	TG100713		+ ICso: 165 nM	+ ICso: 215 nM	+++ IC50: 24 nM	++ IC50: 50 nM			
	VS-5584		++++ ICso: 2.6 nM	+++ ICso: 21 nM	++++IC50: 2.7 nM	++++IC50: 3.0 nM			mTOR
	Taselisib		++++ K: 0.29 nM	+++ K: 9.1 nM	++++K: 0.12 pM	++++K: 0.97 nM	+ ICso: 292 nM	+ ICso: 374 nM	
	CZC24832		10.0.201111	+ ICso: 1.1 µM		+++ ICso: 27 pM	1030. 202 1111	1030.01111	
	IPL-549			1050. T. T pivi		+++ ICm: 16 pM			
	Serabelisib		+++ ICso: 21 pM			1050.10111			
	SEIZUEIISIU		+++ ICso: 24 pM			+ IC as: 159 pM			
	GDC 0226		++ IC50. 34 IIW	+++ K: 26.6 pM	++++K·4 pM	+ IC50. 138 IIW			DINA-PR,BRD4,IIII OR
	GDC-0320		1111 N. 0.2 IIW	N. 20.0 HW	TTTTKI. 4 HW	10.2 mm			
	5AR405				111 IC-1 22.2 mM			++++IC50: 1.2 MM	
	IGR-1202				+++ 1650: 22.2 MM			10 . 45 . 14	
	VPS34 Inhibitor 1							+++ IC ₅₀ : 15 nM	
	GDC-0084		++++ Ki: 2 nM	++ K _i : 46 mM	++++Ki: 3 NM	+++ K _i : 10 hM			miok
	AZD8835		+++ IC ₅₀ : 6.2 nM	++ IC ₅₀ : 431 nM	++++IC50: 5.7 nM	++ IC ₅₀ : 90 nM			
	GSK2269557				++++pKi: 9.9				
	PIK-III		+ IC ₅₀ : 3.96μΜ		+ ICso: 1.2μΜ	+ IC ₅₀ : 3.04μΜ		+++ IC ₅₀ : 0.018µM	ΡΙ4Κβ
	VPS34-IN1							+++ IC50: 25 nM	
	Voxtalisib		++ IC ₅₀ : 39 nM	++ IC ₅₀ : 113 nM	++ IC ₅₀ : 43 nM	+++ IC ₅₀ : 9 nM			DNA-PK,mTOR
	AMG319				+++ IC50: 18 nM	+ IC50: 850 nM			
	AZD8186		++ IC50: 35 nM	++++IC50: 4 nM	+++ IC50: 12 nM				
	PF-4989216		++++ IC50: 2 nM		++++IC50: 1 nM	++ IC ₅₀ : 65 nM		++ IC50: 110 nM	
	Pilaralisib		++ IC ₅₀ : 39 nM	++ IC ₅₀ : 36 nM	++ IC50: 36 nM	+++ IC50: 23 nM			
	PI-3065			+ IC50: 1078 nM	+++ IC50: 15 nM				
	HS-173		++++ IC50: 0.8 nM						
	Quercetin			+ IC ₅₀ : 5.4 μM	+ IC ₅₀ : 3.0 μM	+ IC ₅₀ : 2.4 μM			PKC,Src,Sirtuin
	GSK2636771			1					
	CAY10505					V			
	LY3023414	V							DNA-PK,mTOR kinase
	GSK2292767				1				
	GNE-317	1							

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

BEZ235 (NVP-BEZ235, Dactolisib) is a dual ATP-competitive PI3K and mTOR inhibitor for p110 α /y/ δ / β and mTOR(p70S6K) with IC₅₀ of 4 nM /5 nM /7 nM /75 nM /6 nM in cell-free assays, respectively. BEZ235 inhibits ATR with IC50 of 21 nM in 3T3 TopBP1-ER cell.



S1009 BEZ235 (NVP-BEZ235, Dactolisib)



Data from [Cancer Cell, 2012, 21(2): BEZ235 purchased from Selleck

S1065 Pictilisib (GDC-0941)

Pictilisib (GDC-0941) is a potent inhibitor of PI3K α/δ with IC₅₀ of 3 nM in cell-free assays, with modest selectivity against p110 β (11-fold) and p110y (25-fold). Phase 2.

Size 5 mg 50 mg 200 mg 10 mM/1 mL



S1105 LY294002

LY294002 is the first synthetic molecule known to inhibit PI3Ka/ δ/β with IC50 of 0.5 µM/0.57 µM/0.97 µM in cell-free assays, respectively; more stable in solution than Wortmannin, and also blocks autophagosome formation

Size 10 mg 25 mg 200 mg 10 mM/1 mL BGT226 LY294002 Rapamycin Product Citations (66):

5 15 30 5 15 30 5 15 30 (min) Nature, 2015, 10,1038/nature 14412 p-AKT (S473 Hepatology, 2014, 59(4): 1262-72 p-AKT (T308) -----Data from [Clin Cancer Res, 2011, p-p70S6K (T389) 17(22): 7116-26] a-Actinin LY294002 purchased from Selleck

S2226 Idelalisib (CAL-101, GS-1101)

Idelalisib (CAL-101, GS-1101) is a selective p1105 inhibitor with IC50 of 2.5 nM in cell-free assays, and has been shown to have 40- to 300-fold greater selectivity for $p110\delta$ than for $p110\alpha/\beta/\gamma,$ and 400- to 4000-fold more selectivity for p110 δ than for C2 β , hVPS34, DNA-PK and mTOR.



S2247 Buparlisib (BKM120, NVP-BKM120)

Buparlisib (BKM120, NVP-BKM120) is a selective PI3K inhibitor of p110 α/β/δ/y with IC50 of 52 nM/166 nM/116 nM/262 nM in cell-free assays, respectively. BKM120 has reduced potency against VPS34, mTOR, DNAPK, with little activity towards PI4Kβ. Phase 2.

5 mg 10 mg 50 mg 10 mM/1 mL Size

1



S7018 CZC24832 ΡΙ3Κδ.

5 mg 25 mg

10-fold selectivity over PI3K β and >100-fold selectivity over PI3K α and $\lim_{n \to \infty} p \sum_{i=1}^{n-1} \sum_{i=1}^{n-1} p_{i} \sum_{i=1}^{n-1} p_{i$ Size 10 mg 50 mg

S7016 VS-5584 (SB2343)

S7356 HS-173

Size

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VS-5584 (SB2343) is a potent and selective dual PI3K/mTOR inhibitor for mTOR, PI3Ka/ $\beta/\delta/\gamma$ with IC50 of 3.4 nM and 2.6-21 nM, respectively Phase 1. 10 Size 10 mg 50 mg

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S2638 NU7441 (KU-57788)

NU7441 (KU-57788) is a highly potent and selective DNA-PK inhibitor with IC50 of 14 nM and also inhibits PI3K with IC50 of 5 µM in cell-free assavs.

S1038 PI-103

PI-103 is a multi-targeted PI3K inhibitor for p110 $\alpha/\beta/\delta/\gamma$ with IC₅₀ of 2 nM/3 nM/3 nM/15 nM in cell-free assays, less potent to mTOR/DNA-PK with IC50 of 30 nM/23 nM.



S1169 TGX-221

Size

TGX-221 is a p110 β -specific inhibitor with IC₅₀ of 5 nM in a cell-free assay, 1000-fold more selective for p110β than p110α.



Cancer Cell, 2015, 27(1): 97-108

p110B selective

n1105 selective

TGX-221 purchased from Selleck

S1268 IC-87114

IC-87114 is a selective PI3Kō inhibitor with IC $_{50}$ of 0.5 μM in a cell-free assay, 58-fold more selective for PI3Ko than PI3Ky, and over 100-fold more selective than PI3Kα/β.



p110v selective

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CZC24832 is the first selective PI3Ky inhibitor with IC $_{50}$ of 27 nM, with

HS-173 is a potent PI3Ka inhibitor with IC50 of 0.8 nM.



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PI3K/Akt/mTOR



1CD

p110a selectiv



PI3K / mTOR

S8456 VPS34 inhibitor 1 (Compound 19, PIK-III analogue)	S8330 IPI-549 new
VPS34 inhibitor 1 (Compound 19, PIK-III analogue) is a potent and selective inhibitor of VPS34 with an IC∞ of 15 nM.	IPI-549 is a potent inhibitor of PI3K- γ with >100-fold selectivity over other lipid and protein kinases. The biochemical ICso for PI3K- γ is 16 nM
<u>Size 2 mg 5 mg 25 mg</u>	<u>Size 5 mg 25 mg 100 mg</u>

mTOR Inhibitors

Inhibitory Selectivity

Inhibitor Name	mTOR	mTORC1	mTORC2	Other
Dactolisib	+++ IC50: 6 nM			p110α,p110γ,p110δ
Rapamycin	++++ IC ₅₀ : ~0.1 nM			
Everolimus	+++ IC50: 1.6 nM-2.4 nM			
AZD8055	++++ IC50: 0.8~0.13 nM			DNA-PK,ΡΙ3Κδ,ΡΙ3Κα
Temsirolimus	+ IC ₅₀ : 1.76 μM			
PI-103	+ ICso: 30 nM			p110α,p110δ,p110β
KU-0063794		++ IC50: ~10 nM	++ IC50: ~10 nM	
Torkinib	++ IC50: 8 nM			p1100,PDGFR,DNA-PK
Ridaforolimus	++++ IC ₅₀ : 0.2 nM			
INK 128	++++ K: 1.4 nM			ΡΙ3Κα,ΡΙ3Κγ,ΡΙ3Κδ
Voxtalisib Analogue	+ IC50: 157 nM			ΡΙ3Κγ,ΡΙ3Κα,ΡΙ3Κδ
Torin 1	+++ IC50: 4.32 nM	+++ IC50: 2 nM	++ ICso: 10 nM	DNA-PK,p110γ,C2α
Omipalisib		++++ K _i : 0.18 nM	++++ K _i : 0.3 nM	p110α,p110δ,p110γ
OSI-027	+++ IC50: 4 nM	+ ICso: 22 nM	+ ICso: 65 nM	ΡΙ3Κγ, DNA-PK, ΡΙ3Κα
PF-04691502	++ K:: 16 nM			ΡΙ3Κδ,ΡΙ3Κα,ΡΙ3Κγ
Apitolisib	+ K _i app: 17 nM			p110α,p110δ,p110γ
GSK1059615	++ IC ₅₀ : 12 nM			ΡΙ3Κα,ΡΙ3Κβ,ΡΙ3Κδ
Gedatolisib	++++ IC50: 1.6 nM			ΡΙ3Κα,ΡΙ3Κγ
WYE-354	+++ IC50: 5 nM			ΡΙ3Κα,ΡΙ3Κγ
Vistusertib	+++ IC50: 2.8 nM			P-Akt (S473),pS6 (S235/236)
Torin 2	++++ IC ₅₀ : 0.25 nM			ATM,ATR,DNA-PK
WYE-125132	++++ IC50: 0.19 nM			
PP121	++ IC50: 13 nM			PDGFR,Hck,VEGFR
WYE-687	+++ IC ₅₀ : 7 nM			ΡΙ3Κα,ΡΙ3Κγ,ρ38α
WAY-600	++ IC ₅₀ : 9 nM			ΡΙ3Κα,ΡΙ3Κγ
ETP-46464	++++ IC50: 0.6 nM			ATR,DNA-PK,ΡΙ3Kα
GDC-0349	+++ Ki: 3.8 nM			ΡΙ3Κα
XL388	++ IC ₅₀ : 9.9 nM	+++ IC50: 8 nM	+ IC ₅₀ : 166 nM	
SF2523	+ IC ₅₀ : 280 nM			DNA-PK,ΡΙ3Κα,ΡΙ3Κγ
CZ415	++ pIC50: 8.07			
CC-223	++ IC50: 16 nM			cFMS,FLT4,DNA-PK
Voxtalisib	+ IC ₅₀ : 157 nM			ΡΙ3Κγ,ΡΙ3Κα,ΡΙ3Κδ
Zotarolimus	+++ IC ₅₀ : 2.8 nM			
Tacrolimus	V			
BGT226	V			ΡΙ3Κα,ΡΙ3Κγ,ΡΙ3Κβ
Palomid 529		\checkmark		
Chrysophanic Acid	*			EGFR

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

S1039	Rapamycin	(Sirolimus)	Licensed by Pfizer	
Rapamvo	cin (Sirolimus)	is a specific	mTOR inhibitor w	ith IC:

₅₀ of ~0.1 nM HEK293 cells.





Rapamycin purchased from Selleck

S1120 Everolimus (RAD001)

Everolimus (RAD001) is an inhibitor of FKBP12 with IC50 of 1.6-2.4 nM in a cell-free assay



4120-31] Everolimus purchased from Selleck

S1555 AZD8055

AZD8055 is a novel ATP-competitive mTOR inhibitor with IC50 of 0.8 nM in MDA-MB-468 cells with excellent selectivity (~1,000-fold) against PI3K isoforms and ATM/DNA-PK. Phase 1.



S1044 Temsirolimus (CCI-779, NSC 683864)

Temsirolimus (CCI-779, NSC 683864) is a specific mTOR inhibitor with IC50 of 1.76 µM in a cell-free assay.

Its effects towards mTOR inhibition and FKBP12 binding is similar to



S1022 Ridaforolimus (Deforolimus, MK-8669)

Size 5 mg 10 mg 50 mg 10 mM/1 mL

Licensed by Pfizer



rapamycin. Phase 3.

Data from [PLoS One, 2013, 8(5): e621041

Product Citations (17):

Autophagy, 2011, 7(2): 176-87

Cancer Res, 2014, 74(14): 3947-58

Temsirolimus purchased from Selleck

nTORC1 selectiv Ridaforolimus (Deforolimus, MK-8669) is a selective mTOR inhibitor AKT --with IC50 of 0.2 nM in HT-1080 cell line; while not classified as a prodrug. 235/236) - -n. se (s

e800701 R-Actin -----

5 mg 50 mg 100 mg 10 mM/1 mL Contra Car BKM120 K11-005229 PI3K/Akt/mTOR - + + + - + + + Product Citations (15): Cell Stem Cell, 2012, 10(2); 210-7 9---- 9---- p-SERF Circ Res, 2010, 107(10): 1265-74 me(hr) 0 1 4 24 0 1 4 24 Data from [Oncogene, 2013, p-IGF18 10.1038/onc.2013.5091 KU-0063794 purchased from Selleck

S2218 Torkinib (PP242)

S1226 KU-0063794

effect on PI3Ks.

Size

mTORC1 selective

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mTORC1 sel

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nTORC1 selective

Torkinib (PP242) is a selective mTOR inhibitor with IC50 of 8 nM in cell-free assays; targets both mTOR complexes with >10- and 100-fold selectivity for mTOR than PI3Kδ or PI3Kα/β/γ, respectively. 5 mg 10 mg 50 mg 10 mM/1 mL Size

KU-0063794 is a potent and highly specific dual-mTOR inhibitor of

mTORC1 and mTORC2 with IC50 of ~10 nM in cell-free assays; no



S7811 MHY1485

MHY1485 is a potent, and cell-permeable mTOR activator, and also potently inhibits autophagy.

Size 10 mg 50 mg 200 mg



S2811 INK 128 (MLN0128)

INK 128 (MLN0128) is a potent and selective mTOR inhibitor with IC50 of 1 nM in cell-free assays; >200-fold less potent to class I PI3K isoforms, superior in blocking mTORC1/2 and sensitive to pro-invasion genes (vs Rapamycin). Phase 1.



S2827 Torin 1

Size

Torin 1 is a potent inhibitor of mTORC1/2 with IC50 of 2 nM/10 nM in cell-free assays; exhibits 1000-fold selectivity for mTOR than PI3K. 10 mg 25 mg 50 mg



Torin 1 purchased from Selleck

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S2624 OSI-027

OSI-027 is a selective and potent dual inhibitor of mTORC1 and mTORC2 with IC50 of 22 nM and 65 nM in cell-free assays, and more than 100-fold selectivity is observed for mTOR than for PI3K α , PI3K β , PI3Ky or DNA-PK. Phase 1. NHS CAH

Size 5 mg 10 mg 50 mg 10 mM/1 mL

PI3K/Akt/mTOR

S2783 Vistusertib (AZD2014)

AZD2014 is a novel mTOR inhibitor with IC50 of 2.8 nM in a cell-free assay; highly selective against multiple PI3K isoforms (α/β/γ/δ). AZD2014 showed no or weak binding to the majority of kinases when tested at 1 uM.

Size 5 mg 10 mg



S2817

respectively.

Torin 2

Size 5 mg 10 mg 50 mg 10 mM/1 mL

Torin 2 is a potent and selective mTOR inhibitor with IC50 of 0.25 nM in

p53-/- MEFs cell line; 800-fold greater selectivity for mTOR than PI3K

and improved pharmacokinetic properties; inhibition of

ATM/ATR/DNA-PK with EC50 of 28 nM/35 nM/118 nM, in PC3 cell lines

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Akt Inhibitors

Inhibitory Selectivity

Inhibitor Name	Akt	Akt1	Akt2	Akt3	Other
MK-2206 2HCI		+++ IC50: 8 nM	+++ IC50: 12 nM	+ IC ₅₀ : 65 nM	
Perifosine	+ IC ₅₀ : 4.7 μM				
GSK690693		++++ IC50: 2 nM	+++ IC50: 13 nM	+++ IC50: 9 nM	PKCθ,PKCη,PrkX
Ipatasertib		++++ IC50: 5 nM	++ IC50: 18 nM	+++ IC50: 8 nM	
AZD5363		++++ IC50: 3 nM	+++ IC50: 8 nM	+++ IC ₅₀ : 8 nM	ROCK2
PF-04691502	++++ IC50: 3.8~7.5 nM				ΡΙ3Κδ,ΡΙ3Κα,ΡΙ3Κγ
AT7867		++ IC50: 32 nM	+++ IC50: 17 nM	++ IC50: 47 nM	PKA,p70 S6K
Triciribine	+ IC ₅₀ : 130 nM				HIV-1
CCT128930			++++ IC50: 6 nM		p70 S6K,PKA
A-674563		+++ Ki: 11 nM			PKA,CDK2,GSK-3β
PHT-427	+ Κ.: 2.7 μM				PDK-1
Akti-1/2		++ IC ₅₀ : 58 nM	+ IC ₅₀ : 210 nM	+ IC ₅₀ : 2119 nM	
Uprosertib		+ IC ₅₀ : 180 nM	+ IC ₅₀ : 328 nM	++ IC50: 38 nM	
Afuresertib		++++ Ki: 0.08 nM	++++ Ki: 2 nM	++++ Ki: 2.6 nM	
AT13148		++ IC50: 38 nM	+ IC50: 402 nM	++ IC50: 50 nM	PKA,ROCK2,ROCK1
Miltefosine	√				PI3K,PKC
Honokiol	√				MEK
TIC10 Analogue	√				ERK
Deguelin	√				PI3K
TIC10	1				ERK

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com

"+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

S1078 MK-2206 2HCI

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MK-2206 2HCl is a highly selective inhibitor of Akt1/2/3 with IC50 of 8 nM/12 nM/65 nM in cell-free assays, respectively; no inhibitory activities against 250 other protein kinases observed. Phase 2.



S1037 Perifosine (KRX-0401)

Perifosine (KRX-0401) is a novel Akt inhibitor with IC $_{50}$ of 4.7 μM in MM.1S cells, targeting pleckstrin homology domain of Akt. Phase 3.





S2670 A-674563

GSK690693 is a pan-Akt inhibitor targeting Akt1/2/3 with IC50 of 2 nM/13 nM/9 nM in cell-free assays, and is also sensitive to the AGC kinase family: PKA, PrkX and PKC isozymes. Phase 1.

2 mg 5 mg 10 mg 10 mM/1 mL Size the state

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Akt1 selective

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Size

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S2808 Ipatasertib (GDC-0068)

Size 10 mg 50 mg 10 mM/1 mL

Ipatasertib (GDC-0068) is a highly selective pan-Akt inhibitor targeting Akt1/2/3 with IC50 of 5 nM/18 nM/8 nM in cell-free assays, 620-fold selectivity over PKA. Phase 2.

AZD5363 potently inhibits all isoforms of Akt(Akt1/Akt2/Akt3) with IC50

of 3 nM/8 nM/8 nM in cell-free assays, and has similar effect on

Triciribine is a DNA synthesis inhibitor, and also inhibits Akt in PC3 cell

line and HIV-1 in CEM-SS, H9, H9IIIB, U1 cells with IC50 of 130 nM and

20 nM, respectively. Triciribine does not inhibit PI3K/PDK1 and has

5000-fold less activity in cells lacking adenosine kinase. Phase 1/2.

P70S6K/PKA, but lower activity towards ROCK1/2. Phase 2.

Size 5 mg 10 mg 10 mM/1 mL

Size 5 mg 25 mg 10 mM/1 mL

S8019 AZD5363

S1117 Triciribine

akti akti 50oM 100oM S7863 SC79

SC79 is a brain-penetrable Akt phosphorylation activator and an inhibitor towards Akt-PH domain translocation. ,CÔÇ

S7521 Afuresertib (GSK2110183)

Afuresertib (GSK2110183) is a potent, orally bioavailable Akt inhibitor with Ki of 0.08 nM, 2 nM, and 2.6 nM for Akt1, Akt2, and Akt3, respectively. Phase 2.

Size 5 mg

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S7563 AT13148

AT13148 is an oral, ATP-competitive and multi-AGC kinase inhibitor with IC50 of 38 nM/402 nM/50 nM, 8 nM, 3 nM, and 6 nM/4 nM for Akt1/2/3, p70S6K, PKA, and ROCKI/II, respectively. Phase 1.

Size 5 mg 25 mg 100 mg

Size 5 mg 10 mg 50 mg 10 mM/1 mL

S2635 CCT128930

CCT128930 is a potent, ATP-competitive and selective inhibitor of Akt2 with IC50 of 6 nM, 28-fold greater selectivity for Akt2 than for the closely related PKA kinase. a-07XH

<u>Size 5 mg 10 mg 50 mg 10 mM/1 mL</u>

S2310 Honokiol

Honokiol is the active principle of magnolia extract that inhibits Akt-phosphorylation and promotes ERK1/2 phosphorylation. Phase 3. Size 10 mg 25 mg 50 mg 10 mM/1 mL



Product Citation (1): Sensors and Actuators B. 2013, 189: 11-20

Data from [Sensors and Actuators B. 2013, 189: 11-20] onokiol purchased from Selleci

 $\mathrm{Cr}_{h_{1}}^{\mathrm{c}} \mathrm{C}_{h_{2}}^{\mathrm{c}}$

Akt

TLR7 induced cell proliferation is dependent on AKT Product Citations (3): Eur J Pharmacol, 2015, 764: 208-214 Microvasc Res, 2015, 101: 72-81 8000 Data independently produced by Lee lay hoon from National University of Singapore A-674563 purchased from Selleck

A-674563 is an Akt1 inhibitor with K of 11 nM in cell-free assays,

modest potent to PKA and >30-fold selective for Akt1 over PKC.

Size 10 mg 50 mg 200 mg



25 mg	100 mg	

	۵/s	\mathcal{I}_{i}	ζ



5 mg 25 mg 100 mg

S7492 Uprosertib (GSK2141795)

Uprosertib (GSK2141795) is a selective, ATP-competitive, and orally bioavailable Akt inhibitor with IC50 of 180 nM, 328 nM, and 38 nM for Akt 1, 2 and 3, respectively. Phase 2.



PI3K/Akt/mTOR

13K/Akt

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GSK-3 Inhibitors

Inhibitory Selectivity

Inhibitor Name	GSK-3	GSK-3α	GSK-3β	Other
CHIR-99021 HCI		+++ IC50: 10 nM	++++ IC50: 6.7 nM	
SB216763		++ IC50: 34.3 nM	++ IC50: ~34.3 nM	
CHIR-98014		++++ IC ₅₀ : 0.65 nM	++++ IC50: 0.58 nM	
TWS119			++ IC50: 30 nM	
Tideglusib			+ IC50: 60 nM	
SB415286		+ IC ₅₀ : 78 nM	+ IC ₅₀ : ~78 nM	
BIO	++++ IC50: 5 nM			TYK2,CDK5/p35,CDK2/CyclinA
CHIR-99021		+++ IC50: 10 nM	++++ IC50: 6.7 nM	
AZD2858	+ IC50: 68 nM			
AZD1080		+++ IC50: 6.9 nM	++ IC50: 31 nM	
AR-A014418			++ Ki: 38 nM	
TDZD-8			+ ICso: 2 μM	
LY2090314		++++ IC50: 1.5 nM	++++ IC50: 0.9 nM	
BIO-acetoxime		+++ IC ₅₀ : 10 nM	+++ IC ₅₀ : 10 nM	
IM-12			++ IC ₅₀ : 53 nM	
Indirubin			+ ICso: 0.6 μM	CDK2/CyclinA,CDK5/p35,CDK1/CyclinB
Bikinin	\checkmark			
1-Azakenpaullone			1	

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsss) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com

2 "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

S2924 CHIR-99021 (CT99021) HCI

CHIR-99021 HCI (CT99021) is hydrochloride of CHIR-99021, which is TWS119 is a GSK-3β inhibitor with IC₅₀ of 30 nM in a cell-free assay; a GSK-3\alpha/\beta inhibitor with IC_{50} of 10 nM/6.7 nM; CHIR-99021 shows greater than 500-fold selectivity for GSK-3 versus its closest homologs Cdc2 and ERK2.



S1075 SB216763

SB216763 is a potent and selective GSK-3 inhibitor with IC50 of 34.3 nM for GSK-3α and equally effective on inhibiting human GSK-3β.





S7435 AR-A014418 (GSK-36 Inhibitor VIII)

AR-A014418 is an ATP-competitive, and selective GSK3β inhibitor with IC50 and Ki of 104 nM and 38 nM in cell-free assays, without significant inhibition for 26 other kinases tested. w Cult Size 10 mg 50 mg

Size

no GSK control (H1)

(H1)

GSK-3β selective kinases Size 5 mg 25 mg 100 mg





Tideglusib is an irreversible, non ATP-competitive GSK-3β inhibitor with IC50 of 60 nM in a cell-free assay; fails to inhibit kinases with a Cys homologous to Cys-199 located in the active site. Phase 2.



S7198 BIO (GSK-3 Inhibitor IX, 6-bromoindirubin-3-oxime)

BIO is a specific inhibitor of GSK-3 with IC50 of 5 nM for GSK-3α/β in a cell-free assay, showing >16-fold selectivity over CDK5; also a pan-JAK inhibitor.

Size 10 mg 50 mg

S2729 SB415286

SB415286 is a potent GSK3 inhibitor with IC $_{50}/K_{\rm I}$ of 78 nM/31 nM with equally effective inhibition for GSK-3β.

Size 10 mg 50 mg 10 mM/1 mL

S1263 CHIR-99021 (CT99021)

CHIR-99021 (CT99021) is a GSK-3 α and GSK-3 β inhibitor with IC₅₀ of 10 nM and 6.7 nM, respectively. CHIR99201 does not exhibit cross-reactivity against cyclin-dependent kinases (CDKs) and shows a 350-fold selectivity toward GSK-36 compared to CDKs.

Size 2 mg 5 mg 25 mg 100 mg

S7566 IM-12

IM-12 is a selective GSK-3β inhibitor with IC50 of 53 nM, and also enhances canonical Wnt signalling. Size 10 mg 50 mg 200 mg 0.450

ATM/ATR Inhibitors Activator

Inhibitory Selectivity

Inhibitor Name		ATM		ATR	Other
Dactolisib			++++	IC50: 21 nM	p110α,p110γ,mTOR (p70S6K)
KU-55933	++++	IC50: 12.9 nM			
KU-60019	++++	IC50: 6.3 nM			
VE-821			+++	K _i : 13 nM	
Wortmannin	++	IC50: 150 nM			PI3K,DNA-PK,MLCK
Torin 2	+++	EC50: 28 nM	++	EC50: 35 nM	mTOR,DNA-PK
CP-466722	++	IC50: 410 nM			
VE-822	+	IC50: 34 µM	+++	IC ₅₀ : 19 nM	
ETP-46464	+	IC50: 545 nM	+++	IC50: 14 nM	mTOR,DNA-PK,PI3Kα
CGK 733	++	IC50: 200 nM	++	IC50: 200 nM	
AZ20			+++++	IC50: 5 nM	mTOR
AZD6738			++++	IC50: 1 nM	
Schisandrin B			+	IC50: 7.25 µM	

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

www.selleckchem.com

ATM/ATR Inhibitors

S1092 KU-55933 (ATM Kinase Inhibitor)

GSK-38 selective

- A-A

KU-55933 (ATM Kinase Inhibitor) is a potent and specific ATM inhibitor with IC50/Ki of 12.9 nM/2.2 nM in cell-free assays, and is highly selective for ATM as compared to DNA-PK, PI3K/PI4K, ATR and mTOR.



S1570 KU-60019 KU-60019 is an improved analogue of KU-55933, with IC50 of 6.3 nM for ATM in cell-free assays; 270- and 1600-fold more selective for ATM than for DNA-PK and ATR. It is a highly effective radiosensitizer.



S8007 VE-821

VE-821 is a potent and selective ATP competitive inhibitor of ATR with K_{i}/IC_{50} of 13 nM/26 nM in cell-free assays, shows inhibition of H2AX phosphorylation, minimal activity against PIKKs ATM, DNA-PK, mTOR and PI3Ky.



S7102 VE-822

VE-822 is an ATR inhibitor with IC50 of 19 nM in HT29 cells. Size 10 mg 50 mg 3-0-" Ċ Ó

S7050 AZ20

AZ20 is a novel potent and selective inhibitor of ATR kinase with IC50 of 5 nM in a cell-free assay; 8-fold selectivity over mTOR. Size 5 mg 25 mg

S7693 AZD6738

AZD6738 is an orally active, and selective ATR kinase inhibitor with IC50 of 1 nM. Phase 1/2. Size 5 mg 25 mg



Excellent Validation, Technical Support and Prompt Delivery

14





GSK-3ß selective

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TWS119 purchased from Selleck

S2745 CHIR-98014

S1590 TWS119

cell biology

Size

CHIR-98014 is a potent GSK-3α/β inhibitor with IC₅₀ of 0.65 nM/0.58 nM in cell-free assays, with the ability to distinguish GSK-3 from its closest homologs Cdc2 and ERK2.

capable of inducing neuronal differentiation and maybe useful to stem

10 mg 25 mg 50 mg 10 mM/1 mL

The state

Product Citation (1) Stem Cells Dev, 2013, 22(13): 1893-906 Data from [Stem Cells Dev. 2013 22(13): 1893-906] CHIR-98014 purchased from Selleck



5 mg 25 mg 100 mg

LY2090314 is a potent GSK-3 inhibitor for GSK-3a/ß with IC50 of 1.5 nM/0.9 nM; may improve the efficacy of platinum-based chemotherapy regimens, LY2090314 is highly selective towards GSK3 as demonstrated by its fold selectivity relative to a large panel of

ATM/ATR Activator

S4157 Chloroguine Phosphate

Chloroquine Phosphate is a 4-aminoquinoline anti-malarial and anti-rheumatoid agent, also acting as an ATM activator. Size 50 mg

PDK-1 Inhibitors

Inhibitory Selectivity

Inhibitor Name	PDK-1	Other
OSU-03012	++ IC50: 5 μM	
BX-795	++++ IC50: 6 nM	TBK1/IKKɛ,c-Kit,CDK2/CyclinE
BX-912	+++ IC50: 12 nM	PKA,KDR,CDK2/CyclinE
PHT-427	+ Κ.: 5.2 μΜ	Akt

Notes:

1. For more details, such as half maximal inhibitory concentrations (IC $_{\rm SO}s)$ and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

S1106 OSU-03012 (AR-12)

OSU-03012 (AR-12) is a potent inhibitor of recombinant PDK-1 with IC50 of 5 µM in a cell-free assay and 2-fold increasing in potency over OSU-02067



OSU-03012 purchased from Selleck

S6 Kinase Inhibitors

Inhibitory Selectivity

Inhibitor Name	p70 S6K	p70 S6K1		RSK1		RSK2		RSK3		RSK4	Other
BI-D1870			++	IC50: 31 nM	++	IC50: 24 nM	++	IC50: 18 nM	++	IC50: 15 nM	
AT7867	+ IC50: 85 nM										Akt2,PKA,Akt1
PF-4708671		+ IC ₅₀ : 160 nM									
LJI308			+++	IC50: 6 nM	+++	+ IC50: 4 nM	+++	IC50: 13 nM			
LY2584702 Tosylate	++++ IC50: 4 nM										
LY2584702	++++ IC50: 4 nM										
AT13148	+++ IC50: 8 nM		+	IC50: 85 nM							PKA,ROCK2,ROCK1

Excellent Validation, Technical Support and Prompt Delivery

Notes:

16

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com.

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

S1274 BX-795

Size

a levels of C

Relative

Size

0.3

BX-795 is a potent and specific PDK1 inhibitor with IC50 of 6 nM, 140and 1600-fold more selective for PDK1 than PKA and PKC in cell-free assays, respectively. Meanwhile, in comparison to GSK3ß more than 100-fold selectivity observed for PDK1.

10 mg 50 mg 100 mg 10 mM/1 mL Product Citations (4): Proc Natl Acad Sci USA, 2014, 111(49) 17438-43 FEBS J, 2014, 281(17): 3816-27

> Data from [Virology, 2014, 450-451; 182-951 BX-795 purchased from Selleck

S7087 GSK2334470

GSK2334470 is a novel PDK1 inhibitor with IC50 of ~10 nM in a cell-free assay, with no activity for other close related AGC-kinases. Size 10 mg 50 mg

S7517 AZD7545

5 mg 10 mg

AZD7545 is a potent PDHK inhibitor with IC50 of 36.8 nM and 6.4 nM for PDHK1 and PDHK2, respectively. It failed to inhibit PDHK4 at higher concentrations(>10 nM), AZD7545 stimulates PDHK4 activity.





PF-4708671 is a cell-permeable inhibitor of p70 ribosomal S6 kinase

(S6K1 isoform) with K/IC50 of 20 nM/160 nM in cell-free assays;

400-fold greater selectivity for S6K1 than S6K2, and 4- and >20-fold

selectivity for S6K1 than MSK1 and RSK1/2, respectively. First

Product Citations (5):

e903881

Oncotarget, 2014, 5(10): 3145-58

Data from [PLoS One, 2014, 9(2):

Mol Cancer Ther, 2015, 14(3): 799-809

Size 5 mg 10 mg 50 mg 10 mM/1 mL

S2163 PF-4708671 Licensed and M

S6K1-specific inhibitor to be reported.

Size 10 mg 25 mg 10 mM/1 mL

S7704 LY2584702 Tosylate

LY2584702 Tosylate is a selective and ATP-competitive p70S6K inhibitor with IC50 of 4 nM. Phase 1. 0005

Size 10 mg 50 mg

S7698 LY2584702

LY2584702 is a selective and ATP-competitive p70S6K inhibitor with IC50 of 4 nM. Phase 1. Size 5 mg 25 mg 100 mg



Size 50 mg 200 mg

S2542 Phenformin HCI Phenformin HCl is a hydrochloride salt of phenformin that is an anti-diabetic drug from the biguanide class. It activates AMPK,

NH NH



AMPK Inhibitors Activators

Inhibitory Selectivity

Inhibitor Name	АМРК
Dorsomorphin 2HCI	++ Ki: 109 nM
WZ4003	++++ ICso: 20 nM
Dorsomorphin	++ K _i : 109 nM
HTH-01-015	+++ IC ₅₀ : 100 nM

Notes:

1. For more details, such as half maximal inhibitory concentrations (IC50s) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

AMPK Inhibitors

S7306 Dorsomorphin 2HCI

Dorsomorphin 2HCl is a potent, reversible and selective AMPK inhibitor with K of 109 nM in cell-free assays, exhibiting no significant inhibition for several structurally related kinases including ZAPK, SYK, PKC0, PKA, and JAK3. Dorsomorphin 2HCl also inhibits type I BMP receptor activity.

Size 10 mg 50 mg



S7317 WZ4003

WZ4003 is a highly specific NUAK kinase inhibitor with IC50 of 20 nM and 100 nM for NUAK1 and NUAK2 in cell-base assays, respectively, without significant inhibition on 139 other kinases. ççoi.

Size	5 mg	50 mg	.قر

S7840 Dorsomorphin

p70 S6K1 selective

Dorsomorphin is a potent, reversible and selective AMPK inhibitor with K of 109 nM in cell-free assays, exhibiting no significant inhibition for several structurally related kinases including ZAPK, SYK, PKC0, PKA, and JAK3. Dorsomorphin also inhibits type I BMP receptor activity. Size 5 mg 25 mg 100 mg C 0.000

AMPK Activators

5 mg 10 mg 50 mg 10 mM/1 mL

S2697 A-769662

A-769662 is a potent, reversible AMPK activator with EC50 of 0.8 µM, little effect on GPPase/FBPase activity.



Phosphorylation in mouse liver in vivo	Product Citations (4):
GW1516 + +	Cancer Res, 2013, 74(1): 298-308
A-/69662 + +	J Lipid Res, 2014, 55(7): 1254-66
	Data from [J Lipid Res, 2014, 55(7):
AMER	1254-66]
process and a second se	A-769662 purchased from Selleck



Phase 3.

S7898

 $\mathbf{\hat{s}}_{\mathrm{s}}$

Size

AICAR (Acadesine), an AMPK activator, results in accumulation of ZMP, which mimics the stimulating effect of AMP on AMPK and AMPK kinase

increasing activity and phosphorylation.

Size 50 mg 10 mM/1 mL

CL TT IN HO	

S7898	GS	621	new	
GSK62	1 is a s	pecific and potent AM	/IPK activator.	
Size	5 mg	25 mg		0 0
			α.	Y

PI3K/Akt/

TOR







DNA-PK / MELK

PI3K/Akt/mTOR

DNA-PK Inhibitors

Inhibitory Selectivity

Inhibitor Name	DNA-PK	Other
PI-103	++ IC50: 23 nM	p110α,p110δ,p110β
NU7441	+++ IC50: 14 nM	
PIK-75	++++ IC ₅₀ : 2 nM	p110α,p110γ,p110δ
NU7026	+ ICso: 0.23 μM	РІЗК
PP121	+ ICso: 60 nM	PDGFR,Hck,VEGFR
KU-0060648	++++ IC ₅₀ : 8.6 nM	ΡΙ3Κδ,ΡΙ3Κβ,ΡΙ3Κα
Notos:		

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

S2638 NU7441 (KU-57788)

NU7441 (KU-57788) is a highly potent and selective DNA-PK inhibitor with IC50 of 14 nM and also inhibits PI3K with IC50 of 5 µM in cell-free assavs. 90

5 mg 10 mg 50 mg 200 mg Size



Data from [Nucleic Acids Res, 2013, 41(22): 10157-69] NU7441 purchased from Selleck

, ch

S2893 NU7026 (LY293646)

NU7026 is a potent DNA-PK inhibitor with IC50 of 0.23 µM in cell-free assays; 60-fold selective for DNA-PK than PI3K and inactive against both ATM and ATR.



inhibits DNA-PK with IC50 of 2 nM in cell-free assays. S8045 KU-0060648 KU-0060648 is a dual inhibitor of DNA-PK and PI3K α , PI3K β , PI3K δ with IC50 of 8.6 nM and 4 nM, 0.5 nM, 0.1 nM respectively; less inhibition on PI3Ky with IC50 of 0.59 µM. Size 2 mg 25 mg ŝ

PI-103 is a multi-targeted PI3K inhibitor for $p110\alpha/\beta/\delta/\gamma$ with IC50 of 2 nM/3 nM/3 nM/15 nM in cell-free assays, less potent to mTOR/DNA-PK

----- Page 7

PIK-75 is a p110α inhibitor with IC50 of 5.8 nM (200-fold more potently

than p110β), isoform-specific mutants at Ser773, and also potently

S1038 PI-103

S1205 PIK-75

with IC50 of 30 nM/23 nM.



Epigenetics



HDAC Inhibitors

Inhibitory Selectivity

Inhibitor Name	HDAC	HDAC1	HDAC2	HDAC3	HDAC4	HDAC5	HDAC6	HDAC7	HDAC8	HDAC9	HDAC10	HDAC11	HD1	HD2
Vorinostat	++++ ICso: ~10 nM													
Entinostat		++ IC∞: 0.51 µM		+ IC∞: 1.7 µM										
Panobinostat	++++ IC∞: 5~20 nM													
Trichostatin A	++++ ICso: ~1.8 nM													
Mocetinostat		++ IC=0: 0.15 μM	++ IC∞: 0.29 μM	+ IC:0: 1.66 μΜ								+ IC∞: 0.59 μM		
Belinostat	++++ IC=0: 27 nM													
Romidepsin		+++ ICso: 36 nM	+++ ICso: 47 nM											
MC1568												IC	++ ∞: 100 nM~3.4 µ	M
Tubastatin A HCI							+++ IC=0: 15 nM		+ IC:0: 854 nM					
Givinostat													++++ ICso: 7.5~16 nM	+++++ ICso: 10 nM
Dacinostat	+++ IC:0: 32 nM													
CUDC-101	++++ ICso: 4.4 nM	++++ ICso: 4.5 nM	+++ IC=0: 12.6 nM	++++ ICso: 9.1 nM	+++ IC=0: 13.2 nM	+++ ICso: 11.4 nM	++++ ICso: 5.1 nM	++ ICso: 373 nM	++ IC∞: 79.8 nM	++ IC:0: 67.2 nM	+++ IC∞: 26.1 nM			
Quisinostat 2HCI		++++ ICso: 0.11 nM	++++ ICso: 0.33 nM	++++ IC50: 4.86 nM	++++ ICso: 0.64 nM	++++ ICso: 3.69 nM	++ ICso: 76.8 nM	++ ICso: 119 nM	++++ ICso: 4.26 nM	+++ ICso: 32.1 nM	++++ ICso: 0.46 nM	++++ ICso: 0.37 nM		
Pracinostat		+++ ICso: 49 nM	++ ICso: 96 nM	+++ ICso: 43 nM	+++ IC=0: 56 nM	+++ ICso: 47 nM	+ IC∞: 1.008 µM	++ IC=0: 137 nM	++ IC:0: 140 nM	++ ICso: 70 nM	+++ ICso: 40 nM	++ IC=0: 93 nM		

Inhibitory Selectivity

Inhibitor Name	HDAC	HDAC1	HDAC2	HDAC3	HDAC4	HDAC5	HDAC6	HDAC7	HDAC8	HDAC9	HDAC10	HDAC11	HD1	HD2
PCI-34051		+ ICso: 4 μΜ					+ IC∞: 2.9 μM		++++ ICso: 10 nM		+ IC50: 13 μΜ			
Droxinostat				+ ICso: 16.9 μΜ			+ ICso: 2.47 μΜ		+ ICso: 1.46 μΜ					
Abexinostat		+++++ Ki: 7 nM	+++ K: 19 nM	K: 8.2 nM			+++ K: 17 nM		++ IC=0: 280 nM		+++ ICso: 24 nM			
RGFP966				++ IC∞: 80 nM										
AR-42	+++ ICso: 30 nM													
Ricolinostat		++ ICso: 58 nM	+++ IC:0: 48 nM	+++ IC∞: 51 nM	+ ICso: 7 μΜ	+ ICso: 5 μΜ	++++ ICso: 4.7 nM	+ IC∞: 1.4 μM	++ ICso: 100 nM					
Tacedinaline		+ ICso: 0.9 μM	+ ICso: 0.9 μM	+ ICso: 1.2 μM										
CUDC-907		++++ ICso: 1.7 nM	+++++ ICso: 5.0 nM	ICso: 1.8 nM	++ ICso: 409 nM	+ ICso: 674 nM	+++ ICso: 27 nM	++ ICso: 426 nM	++ ICso: 191 nM	++ ICso: 554 nM	++++ ICso: 2.8 nM	++++ ICso: 5.4 nM		
M344	++ ICso: 100 nM													
Tubacin							++++ ICso: 4 nM							
RG2833		+++ K: 32 nM		K: 5 nM										
Resminostat		+++ ICso: 42.5 nM		+++ ICso: 50.1 nM			++ ICso: 71.8 nM							
Tubastatin A							+++ IC=0: 15 nM							
Citarinostat		+++ ICso: 35 nM	+++ IC:0: 45 nM	+++ IC:0: 46 nM			+++++ ICso: 2.6 nM		++ IC=0: 137 nM					
BRD73954			+ ICso: 9 μΜ				+++ ICso: 36 nM		++ ICso: 120 nM					
BG45		+ ICso: 2 μΜ	+ ICso: 2.2 μM	++ ICso: 289 nM										
4SC-202		+ IC∞: 1.20 μM	+ IC50: 1.12 μΜ	+ ICso: 0.57 μΜ										
CAY10603		++ ICso: 271 nM					++++ ICso: 2 pM							
LMK-235					+++ IC:0: 11.9 nM	++++ IC50: 4.2 nM								
Nexturastat A							++++ ICso: 5 nM							
TMP269					++ IC50: 157 nM	++ ICso: 97 nM		+++ ICso: 43 nM		+++ ICso: 23 nM				
HPOB		+ ICso: 2.9 μM	+ ICso: 4.4 μM	+ IC∞: 1.7 μM			++ ICso: 56 nM		+ ICso: 2.8 μM		+ IC₅₀: 3.0 μM			
Valproic acid sodium salt	V													
Scriptaid	4													
Sodium Phenylbutyrate	4													
Tasquinimod					4									

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation. 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

S1030 Panobinostat (LBH589, NVP-LBH589)

Panobinostat (LBH589) is a novel broad-spectrum HDAC inhibitor with Entinostat (MS-275) strongly inhibits HDAC1 and HDAC3 with IC® of IC50 of 5 nM in a cell-free assay. Phase 3.

Size 10 mg 50 mg 200 mg







S1047 Vorinostat (SAHA, MK0683)

Vorinostat (suberoylanilide hydroxamic acid, SAHA) is an HDAC RG2833 (RGFP109) is a brain-penetrant HDAC inhibitor with IC50 of 60 inhibitor with IC50 of ~10 nM in a cell-free assay.



Trichostatin A (TSA) is an HDAC inhibitor with IC50 of ~1.8 nM in

Product Citations (97):

255-65]

Nat Biotechnol, 2015, 10.1038/nbt.3130

Data from [Nat Biotechnol, 2011, 29(3):

SAHA purchased from Selleck

Nat Biotechnol, 2011, 29(3): 255-65



Size

S7292 RG2833 (RGFP109)

S7229 RGFP966 RGFP966 is an HDAC3 inhibitor with IC50 of 0.08 µM in cell-free assay, exhibiting > 200-fold selectivity over other HDAC. Size 10 mg 50 mg

nM and 50 nM for HDAC1 and HDAC3 in cell-free assays, respectively.

S3020 Romidepsin (FK228, Depsipep

Romidepsin (FK228, depsipeptide) is a potent HDAC1 and HDAC2 inhibitor with IC50 of 36 nM and 47 nM in cell-free assays, respectively.



S1484 MC1568

MC1568 is a selective HDAC inhibitor for maize HD1-A with IC50 of 100 nM in a cell-free assay. It is 34-fold more selective for HD1-A than for HD1-B.



S2627 Tubastatin A HCI

Tubastatin A HCl is a potent and selective HDAC6 inhibitor with IC50 of 15 nM in a cell-free assay. It is selective (1000-fold more) against all other isozymes except HDAC8 (57-fold more).



CUDC-101 is a potent multi-target inhibitor against HDAC, EGFR and HER2 with IC50 of 4.4 nM, 2.4 nM, and 15.7 nM, and inhibits class I/II HDACs, but not class III, Sir-type HDACs. Phase 1. 10 mg 50 mg 10 mM/1 mL





Qi....ÇO

cell-free assavs. Size 2 mg 5 mg 10 mg 10 mM/1 mL Jimir MDA-MB-231 BACH1 VA levels ■ 1833 Product Citations (10): Nat Biotechnol, 2015, 10.1038/nbt.3130 Cancer Cell, 2014, 26(4): 534-48 TSA (hrs 0 24 48 MDA-MB-231 1833 0 24 48 Data from [Proc Natl Acad Sci USA, 2014 111(3) E364-731 Actin ____ TSA purchased from Selleck S1085 Belinostat (PXD101) Belinostat (PXD101) is a novel HDAC inhibitor with IC50 of 27 nM in a cell-free assay, with activity demonstrated in cisplatin-resistant tumors. Size 10 mg 100 mg 200 mg 10 mM/1 mL $\mathrm{Opp}_{\mathrm{p}}$ EMT Induction (Snail-1) Product Citations (15): Control Nat Biotechnol, 2011, 29(3): 255-65 Clin Cancer Res, 2014, 10.1158/1078 EN1 Data from [Cell Rep, 2013, 5(6): 1679-89] PXD101 purchased from Selleck S1122 Mocetinostat (MGCD0103, MG0103)

S1045 Trichostatin A (TSA)

Mocetinostat (MGCD0103) is a potent HDAC inhibitor with most potency for HDAC1 with IC50 of 0.15 µM in a cell-free assay, 2- to 10fold selectivity against HDAC2, 3, and 11, and no activity to HDAC4, 5, 6, 7, and 8. Phase 2.



Data from [Blood, 2014, 123(10); Mocetinostat purchased from Selleck

S7324 TMP269

TMP269 is a potent, selective class IIa HDAC inhibitor with IC50 of 157 nM, 97 nM, 43 nM and 23 nM for HDAC4, HDAC5, HDAC7 and HDAC9, respectively. Size 10 mg 50 mg

Excellent Validation, Technical Support and Prompt Delivery



Size

www.selleckchem.com

Givinostat (ITF2357) is a potent HDAC inhibitor for maize HD2, HD1B AR-42 is an HDAC inhibitor with IC50 of 30 nM. Phase 1. and HD1A with IC50 of 10 nM, 7.5 nM and 16 nM in cell-free assays. Size 2 mg 10 mg 50 mg 10 mM/1 mL Phase 2. MO INCLIS 5 mg 10 mg 50 mg 10 mM/1 mL Size

Product Citations (9): PLoS Pathog, 2014, 10(4): e1004071 J Neurosci, 2013, 33(17): 7535-47 STAT1 Data from [J Interferon Cytokine Res, 2014. 10.1089/iir.2014.00221 Givinostat purchased from Selleck

S1096 Quisinostat (JNJ-26481585) 2HCI

Quisinostat (JNJ-26481585) 2HCl is a novel second-generation HDAC inhibitor with highest potency for HDAC1 with $IC_{\rm 50}$ of 0.11 nM in a cell-free assay, modest potent to HDACs 2, 4, 10, and 11; greater than 30-fold selectivity against HDACs 3, 5, 8, and 9 and lowest potency to HDACs 6 and 7. Phase 2.

grand grand 5 mg 10 mg 50 mg 10 mM/1 mL Size □ no treatmen 0.11 nM JNJ 0.22 nM JNJ Product Citations (4): Hoxa1 RARE Nat Commun, 2013, 4: 2735 J Biol Chem, 2014, 289(28): 19519-30 Data from [J Biol Chem, 2014, 289(28): 19519-301 JNJ-26481585 purchased from Selleck

S1515 Pracinostat (SB939)

Pracinostat (SB939) is a potent pan-HDAC inhibitor with IC $_{\rm 50}$ of 40-140 nM with exception for HDAC6. It has no activity against the class III isoenzyme SIRT I. Phase 2.



S2012 PCI-34051

PCI-34051 is a potent and specific HDAC8 inhibitor with IC50 of 10 nM in a cell-free assay. It has greater than 200-fold selectivity over HDAC1 and 6, more than 1000-fold selectivity over HDAC2, 3, and 10.

Size 10 mg 10 mM/1 mL



Data from [J Mol Biol, 2014, pii S0022-2836(14)00131-4] PCI-34051 purchased from Selleck

S8001 Rocilinostat (ACY-1215)

Rocilinostat (ACY-1215) is a selective HDAC6 inhibitor with IC50 of 5 nM in a cell-free assay. It is >10-fold more selective for HDAC6 than HDAC1/2/3 (class I HDACs) with slight activity against HDAC8, minimal activity against HDAC4/5/7/9/11, Sirtuin1, and Sirtuin2. Phase 2.

Size 5 mg 10 mg 50 mg 10 mM/1 mL



Sodium valproate 0 0.001 0.01 0.1 1 10 (µM) Acetylated Hist

S2244 AR-42 (HDAC-42)

Data independently produced by Dr Zhang of Tianiin Medical University Sodium valproate purchased from Selleck

HDAC1 selective

J Neurosci, 2013, 33(17): 7535-47

anon.

S2818 Tacedinaline (CI994)

Tacedinaline (CI994) is an anti-cancer drug which inhibits HDAC1 with IC50 of 0.57 µM in a cell-free assay and causes G1 cell cycle arrest. Phase 3. Size 10 mg 50 mg 10 mM/1 mL

S2759 CUDC-907

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O Dat

CUDC-907 is a dual PI3K and HDAC inhibitor for PI3K and HDAC1/2/3/10 with IC50 of 19 nM and 1.7 nM/5 nM/1.8 nM/2.8 nM, respectively. Phase 1. Size 5 mg 10 mg 10 mM/1 mL

Tubacin is a highly potent and selective, reversible and cell-permeable HDAC6 inhibitor with an IC50 of 4 nM, approximately 350-fold selectivity over HDAC1. Size 5 mg 10 mg



HDAC6 se Tubastatin A is a potent and selective HDAC6 inhibitor with IC50 of 15 nM. It is selective against all the other isozymes (1000-fold) except

HDAC8 (57-fold). ${\rm e}_{\rm r} {\rm I}_{\rm o}$ tesi TBSA MEC17 KD TBSA+MEC17 KD Product Citations (6): Nat Commun. 2014. 5: 3479 -----Cell Rep, 2013, 5(6): 1679-89 -----Data from [Nat Commun, 2014, 5 3479] Tubastatin A (TBSA) purchased from

Selleck

3730		-235		
LMK-	235 is a	selective	inhibitor of HDAC4 a	Ind HDAC5 with IC50 of 11.9
nM ar	nd 4.2 nN	/l, respe	ctively.	
Size	10 ma	50 ma	200 mg	J. J. Marchan

Tasquinimod is an orally active antiangiogenic agent by allosterically

Size 5 mg 25 mg

97500 LMK 335

S7617 Tasquinimod (ABR-215050)

inhibiting HDAC4 signalling. Phase 3.

CAY10603 is a potent and selective HDAC6 inhibitor with IC50 of 2 pM, >200-fold selectivity over other HDACs.

- Citarinostat (ACY-241) is an orally available selective HDAC6 inhibitor with IC50 of 2.6 nM and 46 nM for HDAC6 and HDAC3, respectively. It has 13 to 18-fold selectivity towards HDAC6 in comparison to

S1999 Sodium butyrate

S8464 Citarinostat (ACY-241)

Size 5 mg 25 mg

PARP Inhibitors

Inhibitory Selectivity

-	-			
Inhibitor Name	PARP	PARP1	PARP2	PARP3
Olaparib		+++ IC50: 5 nM	++++ IC50: 1 nM	
Veliparib		++ Ki: 5.2 nM	+++ Ki: 2.9 nM	
Rucaparib	++++ K _i : 1.4 nM			
Talazoparib	++++ IC ₅₀ : 0.58 nM			
AG-14361		+++ Ki: <5 nM		
INO-1001	++ IC50: <50 nM			
A-966492		++++ K _i : 1 nM	++++ Ki: 1.5 nM	
PJ34	+++ EC50: 20 nM			
PJ34 HCI	+++ EC50: 20 nM			
Niraparib		+++ IC50: 3.8 nM	++++ IC50: 2.1 nM	
UPF 1069		+ IC ₅₀ : 8.0 μM	++ IC ₅₀ : 0.3 μΜ	
ME0328		+ IC ₅₀ : 6.3 μM		+ IC ₅₀ : 0.89 μM
NMS-P118		++ Kd: 0.009 μM		
Picolinamide	+ IC50: 95 μM			
Benzamide	+ IC ₅₀ : 3.3 μM			
Niraparib tosylate		+++ IC50: 3.8 nM	+++ IC50: 2.1 nM	
NU1025	+ IC50: 400 nM			
Iniparib		1		
AZD2461	√			
BGP-15	√			

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsss) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

3. Red "V" refers to compounds which do inhibitory effects on the related isoform, but without specific value

Product Citations (37):

ABT-888 purchased from Selleck

41(7): 4080-92]

S1004 Veliparib (ABT-888)

-

Size 10 mg 50 mg 10 mM/1 mL

DMSO PJ-34 ABT888 AZD2281

_ _ _ _ _ _

Veliparib (ABT-888) is a potent inhibitor of PARP1 and PARP2 with K of 5.2 nM and 2.9 nM in cell-free assays, respectively. It is inactive to SIRT2, Phase 3.

S116-APLE

10 mg 50 mg 200 mg 10 mM/1 mL Size Ċ\$~~ Product Citations (7): TE-6 TE-1 Nat Methods, 2013, 10(10): 981-4 • 100 ق € 100 ¢ Cell Metab. 2014. 19(6): 1034-41 Cell Metab. 2014. 19(6): 1034-41 50 50 J Cell Biol. 2014, 206(4): 493-507 0 0.1 1.0 10 0 0.1 1.0 10 100 Data from [Cancer Sci, 2014, 105(2) Data from [Nucleic Acids Res, 2013, Iniparib (uM) Iniparib (µM) 202-101 Iniparib purchased from Sellec

www.selleckchem.com

Ó.,

30 kDa -45 kDa -

33 kDa 53 kDa 45 kDa

Size 50 mg



S2239 Tubacin S8049 Tubastatin A









HDAC1-3.

<u>iii</u>0

S7596 CAY10603

Size 5 mg 25 mg 100 mg

Sodium butyrate, sodium salt of butyric acid, is a histone deacetylase inhibitor and competitively binds to the zinc sites of class I and II histone deacetylases (HDACs). Size 1 g ~^î....

S1087 Iniparib (BSI-201, NSC-746045, IND-71677) Iniparib (BSI-201) is a PARP1 inhibitor with demonstrated effectiveness



PARP / JAK



S7048 Talazoparib (BMN 673)

Talazoparib (BMN 673) is a novel PARP inhibitor with IC50 of 0.58 nM in a cell-free assay. It is also a potent inhibitor of PARP-2, but does not inhibit PARG and is highly sensitive to PTEN mutation. Phase 3. Size 10 mg 50 mg

JAK Inhibitors

Inhibitory Selectivity

Inhibitor Name	JAK1	JAK2	JAK3	Tyk2	Other
Ruxolitinib	+++ IC ₅₀ : 3.3 nM	++++ IC50: 2.8 nM			
Tofacitinib Citrate	++ IC ₅₀ : 112 nM	++ IC ₅₀ : 20 nM	++++ IC50: 1 nM		ROCK2,LCK
AZD1480		++++ IC50: 0.26 nM			
Fedratinib		++++ IC50: 3 nM			FLT3,RET
AT9283		++++ IC ₅₀ : 1.2 nM	++++ IC ₅₀ : 1.1 nM	+++ IC ₅₀ : 1 nM-10 nM	Aurora B,Aurora A,Abl1
AG-490		+ IC ₅₀ : ~10 μM			EGFR,ErbB2
Momelotinib	+++ IC50: 11 nM	+++ IC50: 18 nM	+ ICso: 155 nM		
Tofacitinib	++ IC50: 112 nM	++ IC50: 20 nM	++++ IC50: 1 nM		ROCK2,LCK
WP1066		+ IC ₅₀ : 2.3 μM			STAT3
TG101209		+++ IC ₅₀ : 6 nM	+ IC ₅₀ : 169 nM		RET,FLT3
Gandotinib	++ IC50: 19.8 nM	++++ IC50: 2.52 nM	++ IC50: 48.0 nM	++ IC50: 44 nM	FLT3,FLT4,FGFR2
NVP-BSK805 2HCI	++ IC50: 31.63 nM	++++ IC50: ~0.5 nM	+++ IC50: 18.68 nM	+++ IC50: 10.76 nM	
Baricitinib	+++ IC ₅₀ : 5.9 nM	+++ IC ₅₀ : 5.7 nM		++ IC ₅₀ : 53 nM	
AZ 960		++++ IC ₅₀ : <3 nM			
CEP-33779		++++ IC50: 1.8 nM			
Pacritinib	+ IC ₅₀ : 1.28 μM	++ ICso: 19~23 nM	+ IC50: 520 nM	++ IC50: 50 nM	FLT3 (D835Y),FLT3
WHI-P154			+ IC ₅₀ : 1.8 μM		EGFR,Src,VEGFR

showing 150-fold selectivity for PARP-1 over PARP-2 (Kd 0.009 µM vs

X-0-61.

1.39 µM, respectively).

Size 5 mg 25 mg

Inhibitory Selectivity

Inhibitor Name	JAK1	JAK2	JAK3	Tyk2	Other
XL019	+ IC ₅₀ : 134.3 nM	++++ IC50: 2.2 nM	+ IC ₅₀ : 214.2 nM	+ IC ₅₀ : 348.3 nM	PDGFRβ,FLT3,c-Kit
S-Ruxolitinib	+++ IC50: 3.3 nM	++++ IC50: 2.8 nM	+ IC50: 428 nM	++ IC50: 19 nM	
ZM 39923 HCI	+ pIC50: 4.4		++ pIC50: 7.1		TGM2,EGFR
Decernotinib	+++ IC ₅₀ : 11 nM	+++ K _i : 13 nM	++++ K _i : 2.5 nM	+++ Ki: 13 nM	
Cerdulatinib	+++ IC ₅₀ : 12 nM	+++ IC50: 6 nM	+++ IC ₅₀ : 8 nM	++++ IC50: 0.5 nM	ARK5,MST1,Fms
Filgotinib	+++ IC50: 10 nM	++ IC50: 28 nM	+ ICso: 810 nM	+ IC50: 116 nM	
FLLL32		+ IC50: <5 μM			
BMS-911543		++++ IC50: 1.1 nM	++ IC ₅₀ : 75 nM	++ IC ₅₀ : 66 nM	SET-2
Peficitinib		1			
GLPG0634 analogue		~			
Go6976		1			FLT3,PKCα,PKCβ1
Curcumol		N			

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com . "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

S1378 Ruxolitinib (INCB018424)

Ruxolitinib (INCB018424) is the first potent, selective JAK1/2 inhibitor to enter the clinic with IC50 of 3.3 nM/2.8 nM in cell-free assays, >130-fold selectivity for JAK1/2 versus JAK3. 5 mg 25 mg 100 mg 10 mM/1 mL Size



Cancer Cell, 2015, 28(1): 29-41 Data from [Blood, 2014, 123(24): 3832-42] Ruxolitinib purchased from Selleck

Product Citations (34):

Nat Med, 2015, 10.1038/nm.4013

S5001 Tofacitinib (CP-690550) Citrate

Tofacitinib (CP-690550) Citrate is a novel inhibitor of JAK3 with IC50 of 1 nM in cell-free assays, 20- to 100-fold less potent against JAK2 and JAK1.



S2736 Fedratinib (SAR302503, TG101348)

Fedratinib (SAR302503, TG101348) is a selective inhibitor of JAK2 with IC50 of 3 nM in cell-free assays, 35- and 334-fold more selective for JAK2 versus JAK1 and JAK3. Phase 2.

Size 5 mg 25 mg 50 mg 10 mM/1 mL TG10134



Product Citations (8): PLoS One, 2014, 9(7): e102741 Cancer Res. 2013, 73(20): 6310-22 Data from [J Cell Mol Med, 2013, 17(2)

Product Citations (13):

154(9): 3219-271

Nat Cell Biol. 2015. 17(1): 57-67

Data from [Endocrinology, 2013,

AZD1480 purchased from Selleck

Blood, 2014, 123(10); 1516-24

265-761 AT9283 purchased from Selleck

S7119 Go6976

S1134 AT9283

AT9283 CEP701 TG101209

S2162 AZD1480

Size

Size

100

extent against JAK1. Phase 1.

P-STAT5 -----

STAT5 E E E E

P-ERK _ _ _ _

5 mg 10 mg 50 mg 10 mM/1 mL

- + - + - + GH+E2 - - + + - - AG1478 - - - + + AZD148

2 mg 10 mg 50 mg 10 mM/1 mL

A19283 UEP/0, 1010.....

Jak2

Jak-targeting inhibitor

AZD1480

cell-free assays; also potent to Aurora A/B, Abl(T315I). Phase 2.

TG101348 CVT

AT9283 is a potent JAK2/3 inhibitor with IC50 of 1.2 nM/1.1 nM in

Go6976 is a potent PKC inhibitor with IC50 of 7.9 nM, 2.3 nM, and 6.2 nM for PKC (Rat brain), PKCa, and PKCB1, respectively. Also a potent inhibitor of JAK2 and Flt3. ----- Page 73

AZD1480 is a novel ATP-competitive JAK2 inhibitor with IC50 of 0.26 nM

in a cell-free assay, selectivity against JAK3 and Tyk2, and to a smaller

S8057 Pacritinib (SB1518)

Pacritinib (SB1518) is a potent and selective inhibitor of Janus Kinase 2 (JAK2) and Fms-Like Tyrosine Kinase-3 (FLT3) with IC50 of 23 and 22 nM in cell-free assays, respectively. Phase 3.

5 mg		

S1143 AG-490 (Tyrphostin B42)

AG-490 (Tyrphostin B42) is an inhibitor of EGFR with IC $_{50}$ of 0.1 μM in cell-free assays, 135-fold more selective for EGFR versus ErbB2, also inhibits JAK2 with no activity to Lck, Lyn, Btk, Syk and Src. ----- Page 39

Size

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Ç, O ~n)

JAK2 selective

S2219 Momelotinib (CYT387, LM-1149

Momelotinib (CYT387) is an ATP-competitive inhibitor of JAK1/JAK2 with IC50 of 11 nM/18 nM, ~10-fold selectivity versus JAK3. Phase 3. and an Size

Product Citations (6):

4093-1031

Nat Cell Biol, 2015, 17(1): 57-67

Data from [Blood 2012 120(19)

CYT387 purchased from Selleck

J Clin Invest, 2014, 124(12): 5263-74

10 mg 50 mg 10 mM/1 mL



S2789 Tofacitinib (CP-690550, Tasocitinib)

Tofacitinib (CP-690550, Tasocitinib) is a novel inhibitor of JAK3 with IC50 of 1 nM in cell-free assays, 20- to 100-fold less potent against JAK2 and JAK1

Size	5 mg50 mg	100 mg10 mM	<u>// mL</u>
CP-690550 (µm p-STAT	NK-S1 (-IL-2) soll.) 0 0.5 1.0 2.0 5	KHYG-1 (+IL-2) 0 0.5 1.0 2.0	Product Citations (34): Biochemistry, 2016, 10.1172/JCI81468 Mol Syst Biol, 2015, 11(3): 797
STAT			Data from [Cancer Discov, 2012, 2(7): 591-7] CP-690550 purchased from Selleck

S2796 WP1066

WP1066 is a novel inhibitor of JAK2 and STAT3 with IC50 of 2.30 µM and 2.43 µM in HEL cells; shows activity to JAK2, STAT3, STAT5, and ERK1/2 not JAK1 and JAK3. Phase 1. $\otimes_{\vec{p}_{ij}} \widehat{\mathbb{C}}^*$

Size 10 mg 25 mg 10 mM/1 mL



S2806 CEP-33779

CEP-33779 is a selective JAK2 inhibitor with IC50 of 1.8 nM, >40- and >800-fold versus JAK1 and TYK2. -0-0⁻¹⁰⁻⁰

Size 5 mg 10 mg 10 mM/1 mL

TG101209 is a selective JAK2 inhibitor with IC50 of 6 nM, less potent to Flt3 and RET with IC50 of 25 nM and 17 nM in cell-free assays. ~30-fold selective for JAK2 than JAK3, sensitive to JAK2V617F and MPI W515I /K mutations



S2851 Baricitinib (LY3009104, INCB028050)

Baricitinib (LY3009104, INCB028050) is a selective JAK1 and JAK2 inhibitor with IC50 of 5.9 nM and 5.7 nM in cell-free assays, ~70 and ~10-fold selective versus JAK3 and Tyk2, no inhibition to c-Met and Chk2 Phase 3 Ď

Size 5 mg 10 mg

S7605 Filgotinib (GLPG064)

5 mg 25 mg 100 mg

Size

Filgotinib (GLPG0634) is a selective JAK1 inhibitor with IC50 of 10 nM, 28 nM, 810 nM, and 116 nM for JAK1, JAK2, JAK3, and TYK2, respectively. Phase 2.



S7634 Cerdulatinib (PRT062070, PRT2070)

Cerdulatinib (PRT-062070) is an oral active, multi-targeted tyrosine kinase inhibitor with IC50 of 12 nM/6 nM/8 nM/0.5 nM and 32 nM for JAK1/JAK2/JAK3/TYK2 and Syk, respectively. Also inhibits 19 other tested kinases with IC50 less than 200 nM. Size 10 mg 50 mg 200 mg



Pim Inhibitors

Inhibitory Selectivity

			-				
nhibitor Name		Pim1		Pim2		Pim3	Other
GI-1776 free base	++	IC50: 7 nM	÷	IC50: 363 nM	+	IC50: 69 nM	FLT3
SMI-4a	++	IC50: 17 nM					
PIM447	++++	K _i : 6 pM	++++	K _i : 18 pM	++++	K _i : 9 pM	
CX-6258 HCI	+++	IC50: 5 nM	+	IC50: 25 nM	++	IC50: 16 nM	

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

S2198 SGI-1776 free base

Notes:

Pim1 selective SGI-1776 free base is a novel ATP competitive inhibitor of Pim1 with IC50 of 7 nM in a cell-free assay, 50- and 10-fold selective versus Pim2 and Pim3, also potent to Flt3 and haspin. Phase 1. 5 mg 10 mg 50 mg 10 mM/1 mL Size



3992-40001 SGI-1776 free base purchased from Selleck

Pim1 selective

S8005 SMI-4a (TCS PIM-1 4a)

SMI-4a is a potent inhibitor of Pim1 with IC50 of 17 nM, modestly potent to Pim-2, and does not significantly inhibit any other serine/threonine- or tyrosine-kinases. Size 10 mg 50 mg 10 mM/1 mL

S7104 AZD1208

NN NA

AZD1208 is a potent, and orally available Pim kinase inhibitor with IC50 of 0.4 nM, 5 nM, and 1.9 nM for Pim1, Pim2, and Pim3 in cell-free assays, respectively. Phase 1 10 mg 50 mg 200 mg Size

HIF Inhibitors Inhibitory Selectivity

			-		
Inhibitor Name		HIF		HIF1	Other
KC7F2	+	IC50: 20 µM	+	IC50: 20 µM	
Roxadustat	1				
2-Methoxyestradiol	1				Microtubule Associated
PX-478 2HCI	1				
BAY 87-2243	1				

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without

S1007 Roxadustat (FG-4592)

Roxadustat (FG-4592) is an HIF- α prolyl hydroxylase inhibitor in a cell-free assay, stabilizes HIF-2 and induces EPO production. Phase 3. Size 10 mg 10 mM/1 mL

S1233 2-Methoxyestradiol (2-MeOE2)

2-Methoxyestradiol (2-MeOE2) depolymerizes microtubules and blocks HIF-1a nuclear accumulation and HIF-transcriptional activity. Ph Size 10 mg 50 mg 100 mg 10 mM/1 mL

Aurora Kinase Inhibitors



lase 2.	







Size

Size

Inhibitory Selectivity Inhibitor Name Aurora A Aurora B Aurora C Other Alisertib ++++ IC50: 1.2 nM Tozasertib ++++ Ki app: 0.6 nM ++ K_i app: 18 nM +++ Ki app: 4.6 nM Bcr-Abl FLT3 ++++ IC50: 0.37 nM Barasertib ZM 447439 + ICso: 130 nM LCK Src MEK1 + ICso⁺ 110 nM MLN8054 +++ IC50: 4 nM IC50: 172 nM LCK,PKA,CK2 Danusertib +++ IC50: 13 nM ++ IC50: 79 nM ++ IC50: 61 nM Abl,TrkA,RET AT9283 ++++ ICso: ~3.0 nM ++++ IC50: ~3.0 nM JAK3, JAK2, Abl1 (T315I) JNJ-7706621 CDK2/CyclinE,CDK2/CyclinA,CDK1/CyclinB +++ IC50: 11 nM +++ IC50: 15 nM + IC₅₀: 250 nM Hesperadin TbAUK1 Aurora A Inhibitor I ++++ IC50: 3.4 nM + IC₅₀; 3.4 μM + IC50: 432 nM KW-2449 ++ ICso: 48 nM FLT3 (D835Y),Abl (T315I),FLT3 SNS-314 Mesylate +++ IC50: 9 nM ++ IC50: 31 nM ++++ IC50: 3 nM ENMD-2076 +++ IC50: 14 nM IC50: 350 nM FLT3,RET,VEGFR3/FLT4 PHA-680632 ++ IC50: 27 nM + IC50: 135 nM + IC50: 120 nM EGER1 PLK1 ELT3 MK-5108 ++++ IC50: 0.064 nM CYC116 +++ K: 8 nM +++ K: 9 nM VEGFR2,FLT3,CDK2/CyclinE AMG-900 +++ IC50: 5 nM ++++ IC50: 4 nM ++++ IC50: 1 nM p38α,TYK2,JNK2 PE-03814735 +++ ICso: 5 nM ++++ ICso: 0.8 nM FLT1.FAK.TrkA CCT129202 ++ IC50: 42 nM + IC50: 198 nM + IC50: 227 nM GSK1070916 ++++ IC50: 3.5 nM +++ IC50: 6.5 nM FLT1,Tie-2,SIK ++ IC₅₀: 15 nM TAK-901 ++ IC50: 21 nM JAK3.c-Src.YES1

Notes: 1. For more details, such as half maximal inhibitory concentrations (ICsss) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com

HIF / Aurora Kinase

S2919 IOX2

IOX2 is a potent inhibitor of HIF-1a prolyl hydroxylase-2 (PHD2) with IC50 of 21 nM in a cell-free assay, >100-fold selectivity over JMJD2A, JMJD2C, JMJD2E, JMJD3, or the 2OG oxygenase FIH.

Size 10 mg 50 mg 10 mM/1 mL ಹಿಗ Ô

S7309 BAY 87-2243

S7612 PX-478 2HCI

factor-1α (HIF-1α) inhibitor. Phase 1

5 mg 25 mg 100 mg

BAY 87-2243 is a potent and selective hypoxia-inducible factor-1 (HIF-1) inhibitor, Phase 1. Size 10 mg 50 mg Ś. St 8-07

PX-478 2HCl is an orally active, and selective hypoxia-inducible

concentrations of each inhibitor, please visit the website of www.selleckchem.co 2 "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation specific value.

S8443 MK-8617 MK-8617 is an orally active pan-inhibitor of Hypoxia-inducible factor prolyl hydroxylase 1-3 (HIF PHD1-3), inhibiting PHD1, 2, 3 with IC50s of 1.0, 1.0 and 14 nM, respectively. 5 mg 25 mg , Sta

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

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Aurora Kinase

S1133 Alisertib (MLN8237) S1100 MLN8054 MLN8054 is a potent and selective inhibitor of Aurora A with IC $_{50}$ of 4 nM Alisertib (MLN8237) is a selective Aurora A inhibitor with IC50 of 1.2 nM in a cell-free assay. It has >200-fold higher selectivity for Aurora A than in Sf9 insect cell. It is more than 40-fold selective for Aurora A than Aurora B. Phase 3. Aurora B. Phase 1. Size 5 mg 10 mg 50 mg 10 mM/1 mL , SCHD Size 5 mg 10 mg 50 mg 10 mM/1 mL ō uM MI N8054 0-01 Product Citations (45); ° ° ° ° ° ° ° ° Cell Stem Cell, 2012, 11(2): 179-94 BimEL Nat Cell Biol, 2015, 17(2): 113-22 Product Citations (10): Cancer Discov, 2014, 4(11): 1281-9 P-Ser 93/94/98 BimEL J Cell Biol, 2012, 198(4): 591-605 Data from [Nat Commun, 2013, 4: Cdc27 26561 MLN8237 purchased from Selleck Data from [Cell Death Differ, 2013, B-Actir 20(10): 1393-403] 1 2 3 4 5 6 7 8 9 S1048 Tozasertib (VX-680, MK-0457) MLN8054 purchased from Selleck Tozasertib (VX-680, MK-0457) is a pan-Aurora inhibitor, mostly against S1107 Danusertib (PHA-739358) Aurora A with Kiapp of 0.6 nM in a cell-free assay, less potent towards Aurora B/Aurora C and 100-fold more selective for Aurora A than 55 Danusertib (PHA-739358) is an Aurora kinase inhibitor for Aurora A/B/C

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CDK2 etc. Phase 2.

DMSO

S1249 JNJ-7706621

S1529 Hesperadin

S1451 Aurora A Inhibitor I

Aurora A than Aurora B.

vivo.

Size

has no activity on Plk1 and Wee1.

Size

Size 25 mg 100 mg 250 mg 10 mM/1 mL VX-680 + MG132 treatment

other kinases Phase 2



S1147 Barasertib (AZD1152-HQPA)

Barasertib (AZD1152-HQPA) is a highly selective Aurora B inhibitor with IC50 of 0.37 nM in a cell-free assay, ~3700 fold more selective for Aurora B over Aurora A. Phase 1.



S1103 ZM 447439

ZM 447439 is a selective and ATP-competitive inhibitor for Aurora A and Aurora B with ICso of 110 nM and 130 nM, respectively. It is more than 8-fold selective for Aurora A/B than MEK1, Src, Lck and has little effect against CDK1/2/4, Plk1, Chk1, etc.





S1134 AT9283

AT9283 is a potent JAK2/3 inhibitor with IC50 of 1.2 nM/1.1 nM in cell-free assays; also potent to Aurora A/B, Abl(T315I). Phase 2. ----- Page 25



G₂-M

Aurora A Inhibitor I is a novel, potent and selective inhibitor of Aurora A

with IC50 of 3.4 nM in a cell-free assay. It is 1000-fold more selective for

with IC50 of 13 nM/79 nM/61 nM in cell-free assays, modestly potent to

Abl, TrkA, c-RET and FGFR1, and less potent to Lck, VEGFR2/3, c-Kit,

LDE-225 + PHA-739358

JNJ-7706621 is pan-CDK inhibitor with the highest potency on CDK1/2

with IC50 of 9 nM/4 nM, showing >6-fold selectivity for CDK1/2 than CDK3/4/6 in cell-free assays. It also potently inhibits Aurora A/B and

Page 80

Hesperadin potently inhibits Aurora B with IC50 of 250 nM in a cell-free

assay. It markedly reduces the activity of AMPK, Lck, MKK1, MAPKAP-K1, CHK1 and PHK while it does not inhibit MKK1 activity in

Product Citations (11): Cancer Res. 2014. 74(20): 5878-90

Product Citations (11):

2825-341

Nature, 2015, 522(7557); 492-6

Nat Cell Biol. 2014. 16(12): 1257-64

Data from [Cancer Res, 2014, 74(10):

Hesperadin purchased from Selleck

6310-22]

Cancer Res, 2013, 73(20): 6310-22

Data from [Cancer Res, 2013, 73(20):

PHA-739358 purchased from Selleck

Xal-

Aurora A selectiv

5 mg 10 mg 50 mg 10 mM/1 mL

DMB-012

LDE-225 PHA-739358

5 mg 10 mg 50 mg 10 mM/1 mL



0.064 nM in a cell-free assay and is 220- and 190-fold more selective for Aurora A than Aurora B/C, while it inhibits TrkA with less than 100-fold selectivity. Phase 1. ${}^{\circ}\dot{U}^{\circ}O_{c}^{*}$

Size 5 mg 10 mg 10 mM/1 mL



S2719 AMG-900 a,d AMG-900 is a potent and highly selective pan-Aurora kinases inhibitor for Aurora A/B/C with IC50 of 5 nM/4 nM /1 nM. It is >10-fold selective for Aurora kinases than p38a, Tyk2, JNK2, Met and Tie2. Phase 1.

S2158 KW-2449

β, IGF-1R, EGFR. Phase 1.

Size 5 mg 10 mg 50 mg 10 mM/1 mL

S2725 PF-03814735

PF-03814735 is a novel, potent and reversible inhibitor of Aurora A/B with IC50 of 0.8 nM/5 nM, is less potent to FIt3, FAK, TrkA, and minimally active to Met and FGFR1. Phase 1.

KW-2449 is a multiple-target inhibitor, mostly for Flt3 with IC50 of 6.6 nM,

modestly potent to FGFR1, Bcr-Abl and Aurora A; little effect on PDGFR

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Size 5 mg 10 mg 50 mg
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no oc

Sirtuin Inhibitors | Activators

Inhibitory Selectivity

Inhibitor Name	SIRT1	SIRT2	SIRT3	Sirtuin	Other
Selisistat (EX 527)	++++ IC50: 38 nM				
Sirtinol	+ IC ₅₀ : 131 μM	+ IC50: 38 μM			
SirReal2		++++ IC50: 140 nM			
Splitomicin				++ IC ₅₀ : 60 μM	
AGK2		++ IC50: 3.5 μM			
Tenovin-6	++ IC50: 21 μM		+ IC50: 67 μM		p53
Nicotinamide				√	

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsss) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

Sirtuin Inhibitors

S1541 Selisistat (EX 527)

EX527

S2804 Sirtinol

Size 5 mg 10 mg 25 mg 10 mM/1 mL

38 µM in cell-free assays, respectively.

Size 5 mg 25 mg 10 mM/1 mL

Selisistat (EX 527) is a potent and selective SIRT1 inhibitor with IC50 of 38 nM in a cell-free assay, exhibiting >200-fold selectivity against SIRT2 and SIRT3.

Product Citations (8):

298-308]

Sirtinol is a specific SIRT1 and SIRT2 inhibitor with IC50 of 131 µM and

Blood, 2014, 124(1): 121-33

J Pineal Res, 2014, 57(2): 228-38

Data from [Cancer Res, 2014, 74(1):

EX 527 purchased from Selleck



Sirtuin Activators

Size 100 mg 200 mg 500 mg 10 mM/1 mL

S1396 Resveratrol

S7577 AGK2

Resveratrol has a wide spectrum of targets including cyclooxygenases (i.e. COX, IC₅₀<1 µM), lipooxygenases (LOX, IC₅₀=2.7 µM) , kinases, sirtuins and other proteins. It has anti-cancer, anti-inflammatory, blood-sugar-lowering and other beneficial cardiovascular effects.



Product Citations (3): Cell Physiol Biochem, 2015, 35(6) 2255-2271 Molecules, 2014, 19(12); 20570-9

Data independently produced by Dr Johanna Weiss of University Hospita Heidelberg

Resveratrol purchased from Selleck



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SRT1720 is	a selective S	SIRT1 activ	vator with EC ₅₀ of 0.16 µM in	= Epigene
ell-free assa	ay, but is >230	-fold less p	potent for SIRT2 and SIRT3.	[®] Inhibito
Size 5 mg	10 mg50	mg 10 mN	<u>//1 mL</u>	Ĵ
			1 2	- <u>S7110 (+)-JQ1</u>
Sirtinol shSIRT1 #1 - + 130-	- + SRT1720 shSIRT1 #1 130		Product Citations (15):	(+)-JQ1 is a BET BRD4(1/2) in cell
B: α-Elk-1 95- 72-			EMBO J, 2013, 32(6): 791-804	family, but not to
72- IB: α-SIRT1 130- 17- 17- 17- 17- 17- 17- 17- 17	Short 72- exposure 130	Shot exposure	 Data from [EMBO J. 2013. 32(6):	Size 10 mg 2
kDa 1 2	3 4 kDa	1 2 3	791-804]	
			_ SKT1720 purchased from Selleck	S2780 I-BET1
2391 Qu	ercetin (Sopho	pretin)	Sirt1 select	
uercetin, a	natural flavon	oid present	t in vegetables, fruit and wine, is	a BRD3 and BRD4
4-5.4 μM. F	Phase 4.	SINT and		Size 5 mg 1
ize 100 mg			но Quality of the second	ан
	Quercetin			
EGF + Quercetin(µM) 0	+ + + + 0.001 0.01 0.01	+ - 1 0		S1216 PFI-1 (
P-AKT			Data independently produced by Dr	PFI-1 is a high
AKT			Zhang of Tianjin Medical University	inhibitor for BRD4 in a cell-free assa
	*		C Quercetin purchased from Selleck	Size 5 mg 5
7792 SR	T2104 (GSK224	45840)	Sirt2 select	žve
			OIDT4 and a standard based in th	
R12104 (G	SK2245840) i	s a selectiv	ve SIRTT activator involved in th	ie
egulation of	SK2245840) i energy home	s a selectiv ostasis. Ph	lase 2.	ne S7189 I-BET-7
regulation of	SK2245840) i energy home 	s a selectiv ostasis. Ph mg	ase 2.	e S7189 I-BET-7
egulation of	SK2245840) i energy home 	s a selectiv ostasis. Ph Img	ve Sir i 1 activator involved in tri lase 2.	S7189 I-BET-7 I-BET-762 is an cell-free assay, s by macrophages
SR12104 (G regulation of Size 5 mg	SK2245840) i energy home 	s a selectiv ostasis. Ph Img	ve sixt i activator involved in tri lase 2.	 S7189 I-BET-7 I-BET-762 is an cell-free assay, s by macrophages other bromodoma
SR12104 (G egulation of Size 5 mg	SK2245840) i energy home 	s a selectiv ostasis. Ph	ve sixt i activator involved in tri lase 2.	 S7189 I-BET-762 is an cell-free assay, s by macrophages other bromodoma Size 10 mg
SR12104 (G egulation of Size 5 mg	SK2245840) i energy home 25 mg100	s a selectiv ostasis. Ph	ve sixt i activator involved in tri lase 2.	S7189 I-BET-7 BET-762 is an cell-free assay, s by macrophages other bromodoma <u>Size 10 mg</u>
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PFI-3 Notes:

specific value.

CPI-203

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com.

+++ BRD4.IC50: 37 nM

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without



bromodomain inhibitor, with IC50 of 77 nM/33 nM for -free assays, binding to all bromodomains of the BET bromodomains outside the BET family. 25 mg

210151A) is a novel selective BET inhibitor for BRD2, with IC50 of 0.5 μ M, 0.25 μ M, and 0.79 μ M in cell-free elv.

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10 mg	50 mg

PF-6405761)

ly selective BET (bromodomain-containing protein) with IC₅₀ of 0.22 µM and for BRD2 with IC₅₀ of 98 nM iy. 50 mg 10 mM/1 mL Kin-C-N

inhibitor for BET proteins with IC50 of ~35 nM in a uppresses the production of proinflammatory proteins and blocks acute inflammation, highly selective over ain-containing proteins.

talone (RVX-208)

/X-208) is a potent BET bromodomain inhibitor with for BD2 in a cell-free assay, about 170-fold selectivity 2. 20 mg



OTX01 10 to 19	5 is a p 9 nM fo	otent BE r BRD2,	T brom BRD3,	odon and l	nain ir BRD4	nhibito in ce	r with	EC50 assay	rangi /s. Ph	ng fro nase '	om 1.
Size	2 mg	10 mg							÷	534	2



potent CREBBP/EP300 inhibitor with IC50 of 21 nM cell-free assays, respectively. Exhibits 40-fold and 250-fold selectivity for CBP over the first BRD of BRD4 (BRD4(1)) and BRD4(2) respectively. Size 10 mg 50 mg



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S8400 Mivebresib (ABBV-075) Mivebresib(ABBV-075) is a novel BET family bromodomain inhibitor. It

binds bromodomains of BRD2/4/T with similar affinities (Ki of 1-2.2 nM) and highly selective for 18 bromodomain proteins tested (Kd > 1 µM; more than 600-fold selectivity vs. BRD4), but exhibits roughly 10-fold weaker potency towards BRD3 (K of 12.2 nM) and has moderate activity towards CREBBP (K_d = 87 μ M; 54-fold selectivity vs. BRD4). Size 5 mg 25 mg

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new	

20 M M

CPI-0610	is	а	potent	and	selective	benzoisoxazoloazepine	BET
bromodom	nain	inh	nibitor ar	nd cur	rently unde	ergoing human clinical tria	ls for
hematolog	lical	m	alignanc	ies.			N

Size 5 mg 25 mg

S7853 CPI-0610

S8496 EED226

EED226 is a potent, selective, and orally bioavailable a novel allosteric Polycomb repressive complex 2 (PRC2) inhibitor with an IC₅₀ of 23.4 nM when the H3K27me0 peptide was used as substrate and an IC50 of 53.5 nM when the mononucleosome was used as the substrate. It directly binds to the H3K27me3 binding pocket of EED.

Size 5 mg 25 mg

Epigenetic Reader Domain Antagonist

S7088 UNC1215

UNC1215 is a potent and selective MBT (malignant brain tumor) antagonist, which binds to L3MBTL3 with IC50 of 40 nM and KD of 120 nM, 50-fold selective versus other members of the human MBT family.

Size 5 mg 25 mg

Histone Acetyltransferase Inhibitors

Inhibitory Selectivity

Inhibitor Name	Histone Acetyltransferase
C646	+++ Ki: 400 nM
MG149	++ ICso: 74 μM
Remodelin	4
Anacardic Acid	√

Notes:

1. For more details, such as half maximal inhibitory concentrations (IC50s) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation. 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

S7152 C646

C646 is an inhibitor for histone acetyltransferase, and inhibits p300 with a K of 400 nM in a cell-free assay. Preferentially selective for p300 versus other acetvltransferases x2

Size 10 mg 50 mg



MG149 is a potent histone acetyltransferase inhibitor with IC50 of 74 µM and 47 µM for Tip60 and MOF, respectively.

Size 5 mg 25 mg 100 mg



S7582 Anacardic Acid

Anacardic Acid is a potent inhibitor of p300 and p300/CBP-associated factor histone acetyltranferases, which also has antibacterial activity, antimicrobial activity, prostaglandin synthase inhibition, and tyrosinase and lipoxygenase inhibition.

Size 10 mg 50 mg 200 mg

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S1848 Curcumin

Curcumin is the principal curcuminoid of the popular Indian spice turmeric, which is a member of the ginger family (Zingiberaceae). It is an inhibitor of p300 histone acetylatransferase (IC₅₀~25 μ M) and Histone deacetylase; activates Nrf2 pathway and supresses the activation of transcription factor NF-KB. miini

Size 50 mg 10 mM/1 mL

DNA Methyltransferase Inhibitors

Inhibitory Selectivity

Inhibitor Name	DNA Methyltransferase	Other
Decitabine	++++ IC50: 100 ng/mL	
RG108	++ IC ₅₀ : 115 nM	
SGI-1027	+ IC ₅₀ : 8 μM	
Lomeguatrib	+++ ICso: 5 nM	
Azacitidine	1	
Zebularine	A	Cytidine deaminase
Thioguanine	1	
Procainamide HCI	1	Sodium channel

Notes:

1. For more details, such as half maximal inhibitory concentrations (IC50s) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

S1200 Decitabine (Deoxycytidine)

Decitabine is a DNA methyltransferase inhibitor, incorporating into DNA and resulting in hypomethylation of DNA and intra-S-phase arrest of DNA replication. It is used to treat myelodysplastic syndrome (MDS). Size 10 mg 25 mg 100 mg 10 mM/1 mL



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DNA Methyltransferase / Histone Methyltransferase

S1782 Azacitidine

Azacitidine is a nucleoside analogue of cytidine that specifically inhibits DNA methylation by trapping DNA methyltransferases. Size 50 mg 5 g 10 mM/1 mL

S2821 RG108

RG108 is an inhibitor of DNA methyltransferase with IC50 of 115 nM in a cell-free assay, but does not cause trapping of covalent enzymes. Size 10 mg 50 mg 200 mg 10 mM/1 mL



S7113 Zebularine (NSC 309132)

Zebularine is a DNA methylation inhibitor that forms a covalent complex with DNA methyltransferases, and also inhibits cytidinedeaminase with K of 2 µM in a cell-free assay.

Size 10 mg 50 mg 10 mM/1 mL

S7276 SGI-1027 (DNA Methyltransferase Inhibitor II)

SGI-1027 is a DNMT inhibitor with IC50 of 6, 8, 7.5 µM for DNMT1, DNMT3A, and DNMT3B in cell-free assays, respectively.

Size 10 mg 100 mg



SGC 0946 ++++ IC50: 0.3 nM ~ N Entacapone ++ ICso: 151 nM +++ K: 5 nM EP7015666 UNC0379 IC50: 7.9 µM IC50: 13-15 nM EI1 MI-2 MI-3

Inhibitor Name

PFI-2

GSK126

EP70047

BRD477

Inhibitory Selectivity

	+	IC50: 446 nM
	+	IC ₅₀ : 648 nM
	++++	K _i : 0.33 nM
	++	IC50: 9.9 nM
77	+++	ICso: 0.4 nM
D	1	

DNA Methyltransferase

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

S7004 EPZ005687

EPZ005687 is a potent and selective inhibitor of EZH2 with Ki of 24 nM in a cell-free assay, 50-fold selectivity against EZH1 and 500-fold selectivity against 15 other protein methyltransferases. . Kajana 5 mg 25 mg



S7062 Pinometostat (EPZ5676)

EPZ5676 is an S-adenosyl methionine (SAM) competitive inhibitor of protein methyltransferase DOT1L with K of 80 pM in a cell-free assay, demonstrating >37,000-fold selectivity against all other PMTs tested; inhibits H3K79 methylation in tumor. Phase 1. Size 10 mg 50 mg

S7061 GSK126

GSK126 is a potent, highly selective EZH2 methyltransferase inhibitor with IC50 of 9.9 nM, >1000-fold selective for EZH2 over 20 other human methyltransferases.

5 mg 25 mg 100 mg Size

Histone Methyltransferase Inhibitors

Inhibitory Selectivity

Inhibitor Name	DNA Me	ethyltransferase
Pinometostat	++++	Ki: 80 pM
EPZ005687	++	K _i : 24 nM
GSK343	+++	IC50: 4~240 nM
BIX 01294	+	IC50: 2.7 µM
Tazemetostat	+++	IC50: 11 nM
3-deazaneplanocin A HCI	++++	K _i : 50 pM
UNC1999	+++	IC50: 2~45 nM
MM-102	++	IC50: 0.4 µM

S7164 GSK343

GSK343 is a potent and selective EZH2 inhibitor with IC50 of 4 nM in a cell-free assay, showing 60 fold selectivity against EZH1, and >1000fold selectivity against other histone methyltransferases to an Size 5 mg 25 mg



S8006 BIX 01294

BIX 01294 is an inhibitor of G9a histone methyltransferase with IC50 of 2.7 µM in a cell-free assay, reduces H3K9me2 of bulk histones, and also weakly inhibits GLP (primarily H3K9me3); no significant activity observed at other histone methyltransferases. Size 10 mg 25 mg 10 mM/1 mL



14.7 nM. It shows pronounced growth suppressive activity in a panel of

human MLL leukemia cell lines(GI50 at 250 nM-570 nM range), but only

a minimal effect in human leukemia cell lines without MLL

Chaetocin, a natural product from Chaetomium species, is a histone

methyltransferase inhibitor with IC50 of 0.8 µM, 2.5 µM and 3 µM for

dSU(VAR)3-9, mouse G9a and Neurospora crassa DIM5, respectively.

MI-503 is a potent and selective Menin-MLL inhibitor with IC $_{\rm 50}$ of

S7817 MI-503

translocations.

S8068 Chaetocir

1 mg 5 mg

Inhibitors

Size

Size

2 mg 5 mg 25 mg

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and the



2.5 nM and 11 nM in cell-free assays, exhibiting a 35-fold selectivity versus EZH1 and >4, 500-fold selectivity relative to 14 other HMTs. Size 10 mg 50 mg

S7265 MM-102 (HMTase Inhibitor IX)

MM-102 is a high-affinity peptidomimetic MLL1 inhibitor with IC50 of 0.4 µM in a cell-free assay.

Size 2 mg 20 mg

S7165 UNC1999

UNC1999 is a potent, orally bioavailable and selective inhibitor of EZH2 and EZH1 with IC50 of 2 nM and 45 nM in cell-free assays, respectively,

showing >1000-fold selectivity over a broad range of epigenetic and non-epigenetic targets.

Size 5 mg

S7353 EPZ004777

EPZ004777 is a potent, selective DOT1L inhibitor with IC $_{\rm 50}$ of 0.4 nM in a cell-free assay and demonstrates >1,200-fold selectivity for DOT1L over all other tested PMTs.

Size 5 mg 50 mg

S7120 3-Deazaneplanocin A (DZNeP) HCI

3-Deazaneplanocin A (DZNeP) HCl, an analog of adenosine, is a competitive inhibitor of S-adenosylhomocysteine hydrolasewith K of 50 pM in a cell-free assay.

Size 1 mg 5 mg

S7079 SGC 0946

SGC 0946 is a highly potent and selective DOT1L methyltransferase inhibitor with IC50 of 0.3 nM in a cell-free assay, but it is inactive against a panel of 12 PMTs and DNMT1.

Size 10 mg 50 mg

S7294 PFI-2

PFI-2 is a potent, selective, and cell-active lysine methyltransferase SETD7 inhibitor with Ki (app) and IC50 of 0.33 nM and 2 nM, 1000-fold selectivity over other methyltransferases and other non-epigenetic targets.

Size 10 mg 50 mg

S7748 EPZ015666

EPZ015666 is a potent, selective and orally bioavailable PRMT5 inhibitor with K of 5 nM, >20,000-fold selectivity over other PMTs.

Inhibitory Selectivity Inhibito Histone demethylase

Histone Demethylase

Name		·····,····,
GSK J4 HCI	++	JMJD3,IC50: 60 nM
OG-L002	+++	LSD1,IC50: 20 nM
JIB-04	++	JMJD2A,IC50: 445 nM;JMJD2D,IC50: 290 nM;JMJD2E,IC50: 340 nM JMJD3,IC50: 855 nM;JMJD2B,IC50: 435 nM;JARID1A,IC50: 230 nM
CPI-455 HCI	++++	IC50: 10 nM
ORY-1001	+++	LSD1/KDM1A,IC50: 20 nM
GSK J1	+++	JMJD3(KDM6B),IC50: 28 nM;UTX(KDM6A),IC50: 53 nM
GSK-LSD1 2HCI	++++	LSD1,IC50: 16 nM
SP2509	++++	LSD1,IC50: 13 nM
ML324	+	JMJD2,IC50: 920 nM
IOX1	++	KDM2A,IC50: 1.8 µM;KDM5C,IC50: 19 µM;PHD2,IC50: 33 µM;

KDM4E IC50: 2.3 uM:KDM3A IC50: 0.1 uM:KDM6B IC50: 1.6 uM KDM4C IC50: 0.6 µM

Notes:

1. For more details, such as half maximal inhibitory concentrations (IC50S) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

S7070 GSK J4 HCI

GSK J4 HCl is a cell permeable prodrug of GSK J1, which is the first selective inhibitor of the H3K27 histone demethylase JMJD3 and UTX with IC50 of 60 nM in a cell-free assay and inactive against a panel of demethylases of the JMJ family. Size 10 mg 50 mg 10 mM/1 mL



OG-L002 is a potent and specific LSD1 inhibitor with IC50 of 20 nM in a cell-free assay, exhibiting 36- and 69-fold selectivity over MAO-B and

MAO-A, respectively. Size 5 mg 25 mg

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S7816 MI-463

/II-463	is a pot	ent inhit	oitor of M	enin-MLL	interaction	with an I	C50 value	
of 15.3	nM.						5.7	
ize	2 ma	5 ma	25 ma				ČĎ-	

Size 5 mg 25 mg

2 mg 5 mg 25 mg

Size

VEGFR

S7234 IOX1	S7680 SP2509
IOX1 is a potent and broad-spectrum inhibitor of 2OG oxygenases, including the JmjC demethylases.	SP2509 is a selective histone demethylase LSD1 inhibitor with ICso of 13 nM, showing no activity against MAO-A, MAO-B, lactate dehydro -genase and glucose oxidase.
or at	Size 5 mg _ 25 mg _ 100 mg

S7281 JIB-04 (NSC 693627)

JIB-04 is a pan-selective Jumonji histone demethylase	inhibitor w
IC50 of 230, 340, 855, 445, 435, 1100, and 290 nM f	for JARID1
JMJD2E, JMJD3, JMJD2A, JMJD2B, JMJD2C, and JMJD2	2D in cell-fr
assays, respectively.	~
	11

Size 20 mg 50 mg

S7796 GSK2879552 2HCI vith 1A, GSK2879552 2HCl is a potent, selective, orally bioavailable, ree irreversible LSD1 inhibitor with Kiapp of 1.7 µM. Phase 1. $O_{M_{N}^{H}}$ Size 5 mg 25 mg na na na ÷. S8287 CPI-455 HCI CPI-455 is a specific KDM5 inhibitor, elevating global levels of H3K4 trimethylation (H3K4me3) and decreased the number of DTPs in

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S7574 GSK-LSD1 2HCI

GSK-LSD1 2HCl is an irreversible, and selective LSD1 inhibitor with IC₅₀ of 16 nM, > 1000 fold selective over other closely related FAD utilizing enzymes (i.e. LSD2, MAO-A, MAO-B).

Size 5 mg 25 mg 100 mg

multiple cancer cell line models treated with standard chemotherapy or targeted agents. ${\rm exp}^{\rm H} = {\rm exp}^{\rm H} = {\rm exp}^{\rm H}$ Size 5 mg 25 mg





VEGFR Inhibitors

Inhibitory Selectivity

Inhibitor Name	VEGFR1	VEGFR2	VEGFR3	Other
Sorafenib Tosylate		++ IC50: 90 nM		Raf-1,B-Raf,B-Raf (V599E)
Sunitinib Malate		+ IC ₅₀ : 80 nM		Kit,FLT3,PDGFRβ
Cabozantinib	+++ IC50: 12 nM	++++ IC50: 0.035 nM	+++ ICso: 6.0 nM	c-Met,Kit,Axl
Ponatinib		++++ IC50: 1.5 nM		Abl,PDGFRa,FGFR1
Axitinib	++++ IC ₅₀ : 0.1 nM	++++ IC ₅₀ : 0.18-0.2 nM	++++ IC50: 0.1-0.3 nM	PDGFRβ,Kit,PDGFRα
Foretinib	+++ IC ₅₀ : 6.8 nM	++++ IC ₅₀ : 0.86 nM	++++ IC50: 2.8 nM	Met, Tie-2, RON
Vandetanib		++ IC50: 40 nM	+ ICso: 110 nM	
Nintedanib	++ IC50: 34 nM	+++ IC50: 13 nM	+++ IC50: 13 nM	LCK,FLT3,FGFR2
Regorafenib	+++ IC ₅₀ : 13 nM	++++ IC ₅₀ : 4.2 nM	+ IC ₅₀ : 46 nM	RET,Raf-1,Kit
Pazopanib HCI	+++ IC ₅₀ : 10 nM	++ IC ₅₀ : 30 nM	+ IC ₅₀ : 47 nM	FGFR,PDGFR,c-Kit
Cediranib	+++ IC50: 5 nM	++++ IC50: 0.5 nM	++++ IC₅o: ≤3 nM	c-Kit,PDGFRβ,FGFR1
PD173074		+ IC50: 100-200 nM		FGFR1,c-Src

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VEGFR

Inhibitory Selectivity

Inhibitor Name	VEGFR1	VEGFR2	VEGFR3	Other
Dovitinib	+++ IC50: 10 nM	+++ IC ₅₀ : 13 nM	+++ IC50: 8 nM	FLT3,c-Kit,FGFR1
Linifanib	++++ IC50: 3 nM	++++ IC50: 4 nM	+ IC50: 190 nM	CSF-1R,FLT3,Kit
Vatalanib 2HCI	+ IC50: 77 nM	++ IC50: 37-270 nM	+ IC50: 660 nM	PDGFRβ,c-Kit,c-Fms
RAF265		++ EC ₅₀ : 30 nM		B-Raf
Tivozanib	++ IC ₅₀ : 30 nM	+++ IC ₅₀ : 6.5 nM	++ IC ₅₀ : 15 nM	EphB2,PDGFRα,PDGFRβ
Motesanib Diphosphate	++++ IC50: 2 nM	++++ IC50: 3-6 nM	+++ IC50: 6 nM	Kit,RET,PDGFR
Lenvatinib	++ IC50: 22 nM	++++ IC50: 4.0 nM	+++ IC50: 5.2 nM	PDGFRβ,FGFR1,PDGFRα
Orantinib		+ Κ _i : 2.1 μΜ		PDGFRβ,FGFR1
Brivanib	+ IC ₅₀ : 380 nM	++ IC ₅₀ : 25 nM		FGFR1
MGCD-265	++++ IC50: 3 nM	++++ IC50: 3 nM	++++ IC50: 4 nM	Met,RON,Tie-2
AEE788	+ IC50: 59 nM	+ IC50: 77 nM	+ IC50: 330 nM	EGFR,HER2/ErbB2,c-Abl
ENMD-2076		+ IC ₅₀ : 58.2 nM	++ IC ₅₀ : 15.9 nM	FLT3,RET,Aurora A
OSI-930	+++ IC50: 8 nM	+++ IC ₅₀ : 9 nM		CSF-1R,LCK,C-Raf
CYC116		++ Ki: 44 nM		Aurora A, Aurora B, FLT3
Ki8751		++++ IC50: 0.9 nM		c-Kit,PDGFRa,FGFR2
Telatinib		+++ IC ₅₀ : 6 nM	++++ IC50: 4 nM	c-Kit,PDGFRα
Pazopanib	+++ IC ₅₀ : 10 nM	++ IC ₅₀ : 30 nM	+ IC ₅₀ : 47 nM	FGFR,PDGFR,c-Kit
KRN 633	+ IC50: 170 nM	+ IC50: 160 nM	+ ICso: 125 nM	PDGFRα,c-Kit,BTK
SAR131675			++ IC50: 23 nM	
Dovitinib Dilactic Acid	+++ IC ₅₀ : 10 nM	+++ IC ₅₀ : 13 nM	+++ IC50: 8 nM	FLT3,c-Kit,FGFR1
Apatinib		++++ IC50: 1 nM		RET,c-Kit,c-Src
BMS-794833		++ IC50: 15 nM		Met
Cabozantinib malate	+++ IC50: 12 nM	++++ IC50: 0.035 nM	+++ ICso: 6.0 nM	c-Met,Kit,Axl
Brivanib Alaninate	+ IC ₅₀ : 380 nM	++ IC ₅₀ : 25 nM		FGFR1
Golvatinib		++ IC50: 16 nM		c-Met
Semaxanib		+ IC ₅₀ : 1.23 μM		
ZM 323881 HCI		++++ IC50: <2 nM		
ZM 306416	+ IC ₅₀ : 0.33 μM			Src,Abl
ENMD-2076 L-(+)-Tartaric acid		+ IC50: 58.2 nM	++ ICso: 15.9 nM	FLT3,RET,Aurora A
BFH772		++++ IC50: 3 nM		
SU5402		++ IC ₅₀ : 20 nM		FGFR1,PDGFRβ
Sunitinib		+ IC ₅₀ : 80 nM		c-Kit,FLT3 ,Kit
Dovitinib Lactate	+++ IC50: 10 nM	+++ IC50: 13 nM	+++ ICso: 8 nM	FLT3,c-Kit,FGFR1
LY2874455		+++ IC50: 7 nM		FGFR2,FGFR1,FGFR4
SKLB1002		++ IC ₅₀ : 32 nM		
AZD2932		+++ IC50: 8 nM		PDGFRβ,Flt3,c-Kit

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

S1005 Axitinib (AG 013736) Licensed by Pfizer

Axitinib is a multi-target inhibitor of VEGFR1, VEGFR2, VEGFR3, PDGFRβ and c-Kit with IC50 of 0.1 nM, 0.2 nM, 0.1-0.3 nM, 1.6 nM and 1.7 nM in Porcine aorta endothelial cells, respectively.

S1010 Nintedanib (BIBF 1120, Intedanib)

Nintedanib (BIBF 1120) is a potent triple angiokinase inhibitor for VEGFR1/2/3, FGFR1/2/3 and PDGFRα/β with IC₅₀ of 34 nM/13 nM/13 nM, 69 nM/37 nM/108 nM and 59 nM/65 nM in cell-free assays. Phase

S1119 Cabozantinib (XL184, BMS-907351)

Cabozantinib (XL184, BMS-907351) is a potent VEGFR2 inhibitor with IC50 of 0.035 nM and also inhibits c-Met, Ret, Kit, FIt-1/3/4, Tie2, and AXL with IC50 of 1.3 nM, 4 nM, 4.6 nM, 12 nM/11.3 nM/6 nM, 14.3 nM and 7 nM in cell-free assays, respectively.

S1046 Vandetanib (ZD6474)

Vandetanib (ZD6474) is a potent inhibitor of VEGFR2 with IC₅₀ of 40 nM in a cell-free assay. It also inhibits VEGFR3 and EGFR with $\mathsf{IC}_{\scriptscriptstyle 50}$ of 110 nM and 500 nM, respectively. Not sensitive to PDGFRβ, Flt1, Tie-2 and FGFR1 with IC₅₀ of 1.1-3.6 µM. No activity against MEK, CDK2, c-Kit, erbB2, FAK, PDK1, Akt and IGF-1R with IC50 above 10 µM.

Size 25 mg 100 mg 500 mg

Nature, 2015, 10.1038/nature14329 Nat Commun, 2014, 5: 3116 Data from [Nat Commun, 2014, 5: 31161 Vandetanib purchased from Selleck

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Product Citations (16):

S1178 Regorafenib (BAY 73-4506, Fluoro-Sorafenib)

Regorafenib (BAY 73-4506) is a multi-target inhibitor for VEGFR1, VEGFR2, VEGFR3, PDGFRβ, Kit, RET and Raf-1 with IC50 of 13 nM/4.2 nM/46 nM, 22 nM, 7 nM, 1.5 nM and 2.5 nM in cell-free assays, respectively.

Size 5 mg 25 mg 100 mg 10 mM/1 mL

S1035 Pazopanib HCI (GW786034 HCI)

Medium

Pazopanib HCI (GW786034 HCI) is a novel multi-target inhibitor of VEGFR1, VEGFR2, VEGFR3, PDGFR, FGFR, c-Kit and c-Fms with IC_{50} of 10 nM, 30 nM, 47 nM, 84 nM, 74 nM, 140 nM and 146 nM in cell-free assays, respectively. g'u' 10 mg 25 mg 100 mg 10 mM/1 mL Size

Data from [Expert Opin Pharmacother, Pazopanib HCI purchased from Sellect

S1017 Cediranib (AZD2171, NSC-732208

VEGFR2 selective

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Cediranib (AZD2171) is a highly potent VEGFR(KDR) inhibitor with IC50 of <1 nM, and also inhibits FIt1/4 with IC50 of 5 nM/≤3 nM, similar activity against c-Kit and PDGFRβ, 36-, 110-fold and >1000-fold selective more for VEGFR than for PDGFR-α, CSF-1R and Flt3 in HUVEC cells. Phase

S1003 Linifanib (ABT-869, AL39324, RG3635)

Linifanib (ABT-869) is a novel, potent ATP-competitive VEGFR/PDGFR inhibitor for KDR, CSF-1R, FIt-1/3 and PDGFRß with IC50 of 4 nM, 3 nM, 3 nM/4 nM and 66 nM respectively, mostly effective in mutant kinase-dependent cancer cells (i.e. FLT3). Phase 3.

S1207 Tivozanib (AV-951)

Tivozanib (AV-951) is a potent and selective VEGFR inhibitor for VEGFR1/2/3 with IC50 of 30 nM/6.5 nM/15 nM, and also inhibits PDGFR and c-Kit, low activity observed against FGFR-1, Flt3, c-Met, EGFR and IGF-1R. Phase 3.

Size 5 mg 10 mg 50 mg 10 mM/1 mL

Product Citations (6): J Chem Inf Model, 2014, 54(3): 881-93 Cytometry A, 2014, 85(6): 537-47

Data from [Cytometry A, 2014, 85(6): 537-471 Tivozanib purchased from Selleck

S1164 Lenvatinib (E7080)

Lenvatinib (E7080) is a multi-target inhibitor, mostly for VEGFR2(KDR)/VEGFR3(Flt-4) with IC50 of 4 nM/5.2 nM, less potent against VEGFR1/Flt-1, ~10-fold more selective for VEGFR2/3 against FGFR1, PDGFR α/β in cell-free assays. Phase 3.

S2221 Apatinib (YN968D1)

10 mg 50 mg 10 mM/1 mL

Apatinib is an orally bioavailable, selective VEGFR2 inhibitor with IC50 of 1 nM.

Size

VEGFR

VEGFR2 selective

Excellent Validation, Technical Support and Prompt Delivery

S1084 Brivanib (BMS-540215) S2845 Semaxanib (SU5416) Brivanib is an ATP-competitive inhibitor against VEGFR2 with IC50 of 25 Semaxanib (SU5416) is a potent and selective VEGFR(Flk-1/KDR) nM, moderate potency against VEGFR-1 and FGFR-1, but >240-fold inhibitor with IC50 of 1.23 µM, 20-fold more selective for VEGFR than against PDGFR-β. Phase 3. for PDGFRβ, lack of activity against EGFR, InsR and FGFR. Phase 3. - tipe of the offer 5 mg 10 mg 50 mg 10 mM/1 mL 5 mg 10 mg 50 mg 10 mM/1 mL Size Size 120 Product Citations (2): _100 12 80 Genome Biol, 2014, 15(8): 428 S4001 Cabozantinib malate (XL184) VEGFR2 selective Int J Oncol, 2013, 44(3): 959-69 မီ 60 Cabozantinib malate (XL184) is the malate of Cabozantinib, a potent Data independently produced by Dr. MDA-MB-23 VEGFR2 inhibitor with IC50 of 0.035 nM and also inhibits c-Met, Ret, Kit, Yong-Weon Yi from Georgetown Flt-1/3/4, Tie2, and AXL with IC50 of 1.3 nM, 4 nM, 4.6 nM, 12 nM/11.3 University Medical Center 0.01 0.1 10 nM/6 nM, 14.3 nM and 7 nM in cell-free assays, respectively. Brivanib purchased from Selleck 10 mg 50 mg 10 mM/1 mL Size in in S3012 Pazopanib (GW786034) ano Pazopanib is a novel multi-target inhibitor of VEGFR1, VEGFR2, VEGFR3, PDGFR, FGFR, c-Kit and c-Fms with IC50 of 10 nM, 30 nM, Product Citations (12): Cancer Discov, 2014, 4(7): 816-27 47 nM, 84 nM, 74 nM, 140 nM and 146 nM in cell-free assays, Nat Commun, 2014, 5: 3116 respectively. Size 25 mg 100 mg à, a, ó Data from [Cell Death Dis, 2013, 4: e627 XL184 purchased from Selleck Product Citations (6): Cancer Discov, 2013, 3(6): 636-47 Clin Cancer Res, 2014, 19(9): 2368-80 Data from [J Virol, 2010, 85(5): 2296-3031 Pazopanib purchased from Selleck EGFR Inhibitors **Inhibitory Selectivity** Inhibitor Name EGFR/ErbB1 HER2/ErbB2 ErbB3 ErbB4 Other Erlotinib HCI ++++ IC50: 2 nM Gefitinih ++ IC50: 26-57 nM ++ IC₅₀: 10.8 nM +++ IC50: 9.2 nM ICso: 367 nM Lanatinih Ditosvlate Afatinib Neratinib Canertinib Lapatinib AG-490 CP-724714 Dacomitini

Afatinib	++++ IC50: 0.5-10 nM	++ ICso: 14 nM	
Neratinib	+ IC50: 92 nM	+ ICso: 59 nM	KDR,Src
Canertinib	++++ IC50: 1.5 nM	+++ IC ₅₀ : 9.0 nM	
Lapatinib	++ IC ₅₀ : 10.8 nM	++ IC ₅₀ : 9.2 nM + IC ₅₀ : 367 nM	
AG-490	+ IC ₅₀ : 0.1 μM	+ ICso: 13.5 μM	
CP-724714		++ ICso: 10 nM	
Dacomitinib	+++ IC50: 6.0 nM	+ IC ₅₀ : 45.7 nM + IC ₅₀ : 73.7 nM	
WZ4002	++++ IC50: 2-8 nM	+++ IC ₅₀ : 4 nM	
Sapitinib	+++ IC50: 4 nM	+++ ICso: 3 nM	
CUDC-101	+++ IC50: 2.4 nM	++ ICso: 15.7 nM	HDAC,HDAC1,HDAC6
AG-1478	+++ IC ₅₀ : 3 nM		
PD153035 HCI	++++ K _i : 5.2 pM		
Pelitinib	+ IC50: 38.5 nM	+ ICso: 1.255 μM	Src,MEK/ERK,Raf
AEE788	++++ IC50: 2 nM	+++ ICso: 6 nM + ICso: 160 nM	c-Abl,FLT1,c-Fms
AC480	++ IC ₅₀ : 20 nM	+ IC ₅₀ : 30 nM + IC ₅₀ : 190 nM	
OSI-420	++++ IC ₅₀ : 2 nM		
WZ3146	++++ IC50: 2-5 nM		
AST-1306	++++ IC50: 0.5-12 nM	+++ ICso: 3.0 nM ++++ ICso: 0.8 nM	
Rociletinib	++ K _i : 21.5-303.3 nM		
Varlitinib	+++ IC ₅₀ : 7 nM	++++ IC ₅₀ : 2 nM	
Icotinib	+++ IC50: 5 nM		
TAK-285	++ IC50: 23 nM	++ ICso: 17 nM + ICso: 260 nM	MEK1.Aurora B.LCK

Inhibitory Selectivity

Inhibitor Name	EGFR/ErbB1	HER2/ErbB2	ErbB3	ErbB4	Other
WHI-P154	+++ IC50: 4 nM				Src, VEGFR, JAK3
PD168393	++++ IC50: 0.70 nM				
CNX-2006	++ IC50: <20 nM				
Tyrphostin 9	+ IC ₅₀ : 460 μM				PDGFR
AG-18	+ IC ₅₀ : 35 μM				
AZD3759	++++ IC50: 0.3 nM				
Afatinib Dimaleate	++++ IC50: 0.4-0.5 nM	++ IC50: 14 nM			
Erlotinib	++++ IC50: 2 nM				
CL-387785	++++ IC50: 370 pM				
Osimertinib	++ IC50: 12.92 nM				
Genistein	V				topo II
Naquotinib	N				

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com

EGFR/ErbB1 selective

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2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

S1023 Erlotinib HCI (OSI-744)

Erlotinib HCl (OSI-744) is an EGFR inhibitor with IC50 of 2 nM in cell-free assays; >1000-fold more sensitive for EGFR than for human c-Src or

v-Abl. Size 100 mg 500 mg

S1025 Gefitinib (ZD1839)

Gefitinib (ZD1839) is an EGFR inhibitor for Tyr1173, Tyr992, Tyr1173 and Tyr992 in the NR6wtEGFR and NR6W cells with IC50 of 37 nM, 37nM, 26 nM and 57 nM, respectively.

Size 100 mg 250 mg 10 mM/1 mL

S1019 Canertinib (CI-1033, PD183805)

Canertinib (CI-1033) is a pan-ErbB inhibitor for EGFR and ErbB2 with IC50 of 1.5 nM and 9.0 nM; no activity to PDGFR, FGFR, InsR, PKC, or CDK1/2/4. Phase 3.

Size	10 mg	25 mg	50 mg	200 mg	
	<u>EGF</u> (GBN	R-A289D I SKMG3			Product Citations (7):
		HKI-272	[nM] IB: pEGFR (Y1	068)	Cancer Discov, 2012, 2(5): 458-71 PLoS One, 2015, 10(4): e0123623 Data from [Cancer Discov, 2012, 2(5):
			IB: pEGFR (Y1 IB: EGFR	173)	458-71] CI-1033 purchased from Selleck

EGFR/ErbB1 selective S1011 Afatinib (BIBW2992)

> Afatinib (BIBW2992) irreversibly inhibits EGFR/HER2 including EGFR(wt), EGFR(L858R), EGFR(L858R/T790M) and HER2 with IC50 of 0.5 nM, 0.4 nM, 10 nM and 14 nM in cell-free assays, respectively.

S2111 Lapatinib (GW-572016)

Lapatinib, used in the form of Lapatinib Ditosylate, is a potent EGFR and ErbB2 inhibitor with IC50 of 10.8 and 9.2 nM in cell-free assays, respectively.

S1143 AG-490 (Tyrphostin B42)

AG-490 (Tyrphostin B42) is an inhibitor of EGFR with IC50 of 0.1 µM in cell-free assays; 135-fold more selective for EGFR versus ErbB2; also inhibits JAK2 with no activity to Lck, Lyn, Btk, Syk and Src.

EGFR

PDGFR Inhibitors

Inhibitory Selectivity

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Inhibitor Name	PDGFR	PDGFRα	PDGFRβ	Other
Sorafenib Tosylate			++ IC50: 57 nM	Raf-1,VEGFR2/Flk1,B-Raf
Imatinib Mesylate	++ IC50: 100 nM			c-Kit,v-Abl
Sunitinib Malate			++++ IC50: 2 nM	FLT3,Kit,VEGFR2
Ponatinib		++++ IC50: 1.1 nM		Abl,VEGFR2,FGFR1
Axitinib		+++ IC50: 5.0 nM	++++ IC50: 1.6 nM	VEGFR1/FLT1, VEGFR2/Flk1, VEGFR3
Imatinib	++ IC50: 100 nM			c-Kit,v-Abl
Nintedanib		++ IC ₅₀ : 59 nM	++ IC50: 65 nM	VEGFR3,VEGFR2,LCK
Pazopanib HCI	++ IC ₅₀ : 84 nM			VEGFR1,VEGFR2,VEGFR3
Dovitinib		+ IC50: 210 nM	+++ IC50: 27 nM	FLT3,c-Kit,FGFR1
Linifanib			++ IC50: 66 nM	VEGFR1/FLT1,CSF-1R,FLT3
Crenolanib		++++ K _d : 2.1 nM	++++ K _d : 3.2 nM	
Masitinib		+ IC ₅₀ : 540 nM	+ IC50: 800 nM	Kit,Lyn B,Abl1
Tivozanib		+++ IC50: 40 nM	++ IC50: 49 nM	VEGFR2,VEGFR3,EphB2
Amuvatinib		+++ IC50: 40 nM		c-Kit (D816H),FLT3 (D835Y)
Motesanib Diphosphate	++ IC50: 84 nM			VEGFR1,VEGFR2,VEGFR3
Orantinib			+++ Ki: 8 nM	FGFR1,Flk1
CP-673451		+++ IC50: 10 nM	++++ IC50: 1 nM	c-Kit,VEGFR2,VEGFR1
Ki8751		++ IC50: 67 nM		VEGFR2,c-Kit,FGFR2
Telatinib		+++ IC50: 15 nM		c-Kit,VEGFR3,VEGFR2
PP121	++++ IC50: 2 nM			Hck,VEGFR,mTOR
Pazopanib	++ IC50: 84 nM			VEGFR1,VEGFR2,VEGFR3
KRN 633		+ IC50: 965 nM	+ IC ₅₀ : 9850 nM	VEGFR3,VEGFR2,VEGFR1
Dovitinib Dilactic Acid		+ IC ₅₀ : 210 nM	+++ IC50: 27 nM	FLT3,c-Kit,VEGFR3/FLT4
MK-2461			+++ IC50: 22 nM	c-Met (M1250T),c-Met (Y1235D),c-Met (Y1230H)
Tyrphostin AG 1296	+ IC50: 0.3-0.5 μM			c-Kit (Swiss 3T3),FGFR (Swiss 3T3)
Sunitinib			++++ IC ₅₀ : 2 nM	FLT3 ,Kit ,c-Kit
Dovitinib Lactate			+++ IC50: 27 nM	FLT3,c-Kit,VEGFR3/FLT4
AZD2932			+++ IC50: 4 nM	Flt3,VEGFR-2,c-Kit
Sennoside B	V			

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com

PDGFR8 selective

S2475 Imatinib (STI571)

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

S1042 Sunitinib Malate Licensed by Pfizer

Sunitinib Malate is a multi-target RTK inhibitor targeting VEGFR2 Imatinib (STI571) is a multi-target inhibitor of tyrosine kinase with (Flk-1) and PDGFRβ with IC50 of 80 nM and 2 nM in cell-free assays, and also inhibits c-Kit. Size 50 mg 100 mg 500 mg 10 mM/1 mL Size $\mathcal{A}_{\mathcal{D}}$ CHO DMSO + DMSO + Akti1/2 + Akti1/2 Product Citations (41):

	LINSIUND	Nature, 2011, 478(7369): 349-55 Sci Tranel Med, 2015, 7(284): 284r
-		Data from [Leukemia, 2014,
		10.1038/leu.2014.123] Sunitinib Malate purchased from S

S1536 CP-673451

Sunitinit

Actin

d.Caspase9

cl.Caspase

CP-673451 is a selective inhibitor of PDGFRa/ß with IC50 of 10 nM/1 nM in cell-free assays, exhibiting >450-fold selectivity over other angiogenic receptors, and has antiangiogenic and antitumor activity.

Size 5 mg 10 mg 50 mg 10 mM/1 mL

inhibition for v-Abl, c-Kit and PDGFR, IC50 values are 0.6 µM, 0.1 µM and 0.1 µM in cell-free or cell-based assays, respectively. 250 mg 500 mg 10 mM/1 mL Taur ra57 Product Citations (33): Cancer Cell, 2014, 26(6); 840-50 Cell Stem Cell, 2012, 10(2); 210-7 elleck Data from [Blood, 2014, 123(21): 3296-3041

Imatinib purchased from Selleci

Excellent Validation, Technical Support and Prompt Delivery

PDGFR / c-Met

Excellent Validation, Technical Support and Prompt Delivery

Tepotinib (EMD 1214063) is a potent and selective c-Met inhibitor with IC50 of 4 nM, >200-fold selective for c-Met than for IRAK4, TrkA, Axl, IRAK1, and Mer. Phase 1. onan.

Size 5 mg 25 mg

with IC50 of 0.13 nM in a cell-free assay, inactive against RONβ, as well as EGFR and HER-3. Phase 1.

Size

525 min.

<u>1995</u>

Size 5 mg 25 mg

Size 10 mg 50 mg

S8167 AMG 337

inhibitor of the MET receptor.

5 mg 25 mg

S7014 Merestinib (LY2801653)

S2788 Capmatinib (INCB28060)

Capmatinib (INCB28060) is a novel, ATP-competitive inhibitor of c-MET

AMG 337 is an oral, small molecule, ATP-competitive, highly selective

LY2801653 is a type-II ATP competitive, slow-off inhibitor of MET

tyrosine kinase with a dissociation constant (Ki) of 2 nM, a

pharmacodynamic residence time (Koff) of 0.00132 min(-1) and t1/2 of

S1561 BMS-777607

BMS-777607 is a Met-related inhibitor for c-Met, Axl, Ron and Tyro3 with IC50 of 3.9 nM, 1.1 nM, 1.8 nM and 4.3 nM in cell-free assays, 40-fold more selective for Met-related targets versus Lck, VEGFR-2, and TrkA/B, and more than 500-fold greater selectivity versus all other receptor and non-receptor kinases. Phase 1/2.

Size 5 m	g <u>10 mg</u> 50 m	10 mM/1 i	≞ phro,
Control	BMS-777607 +	IS-777607 XAV939	Product Citations (8): J Clin Invest, 2014, 124(11): 4737-52 Mol Cancer Ther, 2014, 13(1): 37-48
1.0 ± 0.1	82.9 ± 7.3 77 % of Poly	.0 ± 7.8 yploid cells	Data from [Mol Cancer Ther, 2014, 13(1): 37-48] BMS-777607 purchased from Selleck

S1114 JNJ-38877605

JNJ-38877605 is an ATP-competitive inhibitor of c-Met with IC₅₀ of 4 nM, 600-fold selective for c-Met than for 200 other tyrosine and serine-threonine kinases. Phase 1.

S2753 Tivantinib (ARQ 197) Tivantinib (ARQ 197) is the first non-ATP-competitive c-Met inhibitor with K of 0.355 μM in a cell-free assay, little activity to Ron, and no inhibition to EGFR, InsR, PDGFRa or FGFR1/4. Phase 3.

10 mg 50 mg Product Citations (2): Cell Signal, 2013, 25(12): 2652-60 PLoS One, 2014, 9(9): e105919 Data from [Cell Signal, 2013, 25(12);

2652-601 3 3 3 3 ARQ 197 purchased from Selleck

S1361 MGCD-265

MGCD-265 is a potent, multi-target and ATP-competitive inhibitor of c-Met and VEGFR1/2/3 with IC50 of 1 nM, 3 nM/3 nM/4 nM, respectively; also inhibits Ron and Tie2. Phase 1/2.

HER2 Inhibitors

Inhibitory Selectivity

Inhibitor Name		HER2	Other
Lapatinib Ditosylate	+++	IC ₅₀ : 9.2 nM	EGFR,ErbB4,c-Src
Afatinib	++	IC50: 14 nM	EGFR (L858R),EGFR (wt),EGFR (L858R/T790M)
Neratinib	+	IC50: 59 nM	EGFR,KDR,Src
Canertinib	+++	IC ₅₀ : 9.0 nM	EGFR
Lapatinib	+++	IC50: 9.2 nM	EGFR,ErbB4,c-Src
CP-724714	+++	IC50: 10 nM	
Sapitinib	++++	IC50: 3 nM	EGFR,ErbB3
CUDC-101	++	IC ₅₀ : 15.7 nM	EGFR,HDAC,HDAC1
Mubritinib	++++	IC50: 6.0 nM	
AEE788	++++	IC50: 6 nM	EGFR,c-Abl,FLT1
AC480	+	IC50: 30 nM	HER1,HER4
TAK-285	++	IC50: 17 nM	EGFR/HER1,HER4,MEK1
Tyrphostin AG 879	+	IC50: 1.0 µM	Trk
Irbinitinib	+++	IC50: 8 nM	p95 HER2
Afatinib Dimaleate	++	IC50: 14 nM	EGFR (L858R),EGFR (wt),EGFR (L858R/T790M)

Notes:

1. For more details, such as half maximal inhibitory concentrations (IC50S) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

S1028 Lapatinib (GW-572016) Ditosylate

Lapatinib (GW-572016) Ditosylate is a potent EGFR and ErbB2 inhibitor with IC50 of 10.8 and 9.2 nM in cell-free assays, respectively. Size

www.selleckchem.com

otein Tvrosine Kinase

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HER2 / IGF-1R

S2150 Neratinib (HKI-272)

Neratinib (HKI-272) is a highly selective HER2 and EGFR inhibitor with IC50 of 59 nM and 92 nM in cell-free assays; weakly inhibits KDR and Src, no significant inhibition to Akt, CDK1/2/4, IKK-2, MK-2, PDK1, c-Raf and c-Met. Phase 3.

6

Data from [Cancer Discov, 2013, 3(2): 224-371

S1167 CP-724714

CP-724714 is a potent, selective inhibitor of HER2/ErbB2 with IC50 of 10 nM, >640-fold selectivity against EGFR, InsR, IRG-1R, PDGFR, VEGFR2, Abl, Src, c-Met etc in cell-free assays. Phase 2.

S2192 Sapitinib (AZD8931)

Sapitinib (AZD8931) is a reversible, ATP competitive inhibitor of EGFR, ErbB2 and ErbB3 with IC50 of 4 nM, 3 nM and 4 nM in cell-free assays, more potent than Gefitinib or Lapatinib against NSCLC cell, 100-fold more selective for the ErbB family than MNK1 and Flt. Phase 2.

S2216 Mubritinib (TAK 165)

Mubritinib (TAK 165) is a potent inhibitor of HER2/ErbB2 with IC50 of 6 nM in BT-474 cell; no activity to EGFR, FGFR, PDGFR, JAK1, Src and Blk in BT-474 cell line. Phase 1.

BM AG-GSł Neratinib purchased from Selleck

IGF-1R Inhibitors Inhibitory Selectivity

Inhibitor Name	IGF-1R	Insulin Receptor	Other
Linsitinib	+++ IC50: 35 nM	++ IC ₅₀ : 75 nM	IRR
NVP-AEW541	++ IC ₅₀ : 0.15 μM	++ IC ₅₀ : 0.14 μM	FLT3,Tek,FLT1
GSK1904529A	+++ IC50: 27 nM	+++ IC50: 25 nM	IKK3,VEGFR2,Syk
NVP-ADW742	+ ICso: 0.17 μM		
BMS-536924	++ IC50: 100 nM	+++ IC ₅₀ : 73 nM	FAK,MEK,LCK
AG-1024	+ IC ₅₀ : 7 μM	+ IC ₅₀ : 57 μM	
GSK1838705A	+++ IC50: 2 nM	++++ IC50: 1.6 nM	ALK,RSK1,JNK3
BMS-754807	++++ IC50: 1.8 nM	++++ IC50: 1.7 nM	TrkB,Met,TrkA
PQ 401	+ IC ₅₀ : <1 μM		
Picropodophyllin	++++ IC50: 1 nM		

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

S1091 Linsitinib (OSI-906)

Linsitinib (OSI-906) is a selective inhibitor of IGF-1R with IC50 of 35 nM in cell-free assays; modestly potent to InsR with IC50 of 75 nM, and no activity towards Abl, ALK, BTK, EGFR, FGFR1/2, PKA etc. Phase 3. Size

S1034 NVP-AEW541 (AEW541)

v-tubulin

NVP-AEW541 is a potent inhibitor of IGF-1R/InsR with IC50 of 150 nM/140 nM in cell-free assays, greater potency and selectivity for IGF-1R in a cell-based assay.

Picropodophyllin (PPP) is a IGF-1R inhibitor with IC50 of 1 nM. It displays selectivity for IGF-1R and does not coinhibit tyrosine phosphorylation the IR, or of a selected panel of receptors less related to IGF-IR(FGF-R, PDGF-R, OR EGF-R).

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Size 5 mg 25 mg 100 mg

S1093 GSK1904529A

GSK1904529A is a selective inhibitor of IGF-1R and IR with IC50 of 27

Size 10 mg 50 mg 10 mM/1 mL

S1088 NVP-ADW742 (GSK 552602A, ADW742)

NVP-ADW742 is an IGF-1R inhibitor with IC50 of 0.17 µM, >16-fold more potent against IGF-1R than InsR; little activity to HER2, PDGFR, VEGFR-2, Bcr-Abl and c-Kit.

Size 5 mg 10 mg 50 mg 10 mM/1 mL

Data from [J Cancer Res Clin Oncol, 2014, 10.1007/s00432-014-1787-z] ADW742 purchased from Selleck

Mol Endocrinol, 2015, 29(3): 373-83

J Cancer Res Clin Oncol, 2014,

10 1007/s00432-014-1787-7

Product Citations (4):

S1234 AG-1024 (Tyrphostin)

AG-1024 (Tyrphostin) inhibits IGF-1R autophosphorylation with IC50 of 7 µM, is less potent to IR with IC50 of 57 µM and specifically distinguishes between InsR and IGF-1R (as compared to other tyrphostins).

5 mg 10 mg 10 mM/1 mL Size

AG1204	0	5	10	15	
(Mu)			_	F -	Raf1
[-	pSrc family (Tyr416)
	=	-	-	4	pAkt (Ser473)
İ				-	Akt
1		-	-	10-s	pMEK (Ser217/221)
[-	-	-	-	MEK
[-	-	-		pERK (Thr202/204)
[ERK
		-	-	-	β-Actin

Data from [Blood, 2013, 122(9): 1621-331 AG-1024 purchased from Selleck

Product Citations (5):

Sci Rep, 2015, 5: 11634

Blood, 2013, 122(9): 1621-33

S1124 BMS-754807

BMS-754807 is a potent and reversible inhibitor of IGF-1R/InsR with IC50 of 1.8 nM/1.7 nM in cell-free assays. less potent to Met. Aurora A/B. TrkA/B and Ron, and shows little activity to Flt3, Lck, MK2, PKA, PKC etc. Phase 2

Size 5 mg 10 mg 50 mg

Product Citations (4): Cancer Res, 2014, 74(20): 5866-77 Clin Cancer Res, 2013, 19(11): 2984-94

Data from [Clin Cancer Res, 2013, 19(11): 2984-941 BMS-754807 purchased from Selleck

www.selleckchem.com

FLT3 Inhibitors

Inhibitory Selectivity

Inhibitor Name	Insu	lin Receptor	Other
Quizartinib	+++	IC50: 1.1-4.2 nM	
Dovitinib	++++	IC50: 1 nM	c-Kit,VEGFR3/FLT4,FGFR1
Amuvatinib	+	IC50: 81 nM	c-Kit (D816H),PDGFRa (V561D)
KW-2449	++++	IC50: 1-6.6 nM	Abl (T315I),Abl,FGFR1
Dovitinib Dilactic Acid	++++	IC50: 1 nM	c-Kit,FGFR1,VEGFR3/FLT4
ENMD-2076 L-(+)-Tartaric acid	++	IC50: 1.86 nM	RET, Aurora A, VEGFR3/FLT4
UNC2025	++++	IC ₅₀ : 0.8 nM	Mer,Axl,Tyro3
G-749	++++	IC50: 0.4-0.6 nM	Mer,Aurora B,RET
AZD2932	++	IC50: 7 nM	PDGFRβ,VEGFR-2,c-Kit
R406	1		Syk
Go6976	1		JAK2,PKCα,PKCβ1

G Notes:

Size

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1. For more details, such as half maximal inhibitory concentrations (IC $_{\rm SO}s)$ and working concentrations of each inhibitor, please visit the website of www.selleckchem.com.

Quizartinib (AC220) is a second-generation FLT3 inhibitor for

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2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

S. 3. Red ">" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

β, RET, and CSF-1R. Phase 3.

2826-371

AC220 purchased from Sellec

S1018 Dovitinib (TKI-258, CHIR-258)

Dovitinib (TKI258, CHIR258) is a multitargeted RTK inhibitor, mostly for class III (FLT3/c-Kit) with IC50 of 1 nM/2 nM, also potent to class IV (FGFR1/3) and class V (VEGFR1-4) RTKs with IC50 of 8-13 nM, less potent to InsR, EGFR, c-Met, EphA2, Tie2, IGF-1R and HER2 in cell-free assays. Phase 4.

S2158 KW-2449

KW-2449 is a multiple-target inhibitor, mostly for Flt3 with IC50 of 6.6 nM, modestly potent to FGFR1, Bcr-Abl and Aurora A; little effect on PDGFR B. IGF-1R. EGFR. Phase 1.

Size 1	10 mg		50 mg	10	mM/1 n	nL		ö0-5
KW2449 (J	JM)	0	0.001	0.01	0.1	1	10	β. H
p-Histon		-	-	-				Data independently produced by
Histon	e -	-	-	-	-	-	-	Dr. Zhang of Tianjin Medical University KW-2449 purchased from Selleck

nM and 25 nM in cell-free assays, >100-fold more selective for IGF-1R/InsR than Akt1/2, Aurora A/B, B-Raf, CDK2, EGFR etc.

FGFR Inhibitors

Inhibitory Selectivity

Inhibitor Name	FGFR	FGFR1	FGFR2	FGFR3	FGFR4	Other
Ponatinib		++++ IC50: 2.2 nM				Abl,PDGFRa,VEGFR2
BGJ398		++++ IC50: 0.9 nM	++++ IC50: 1.4 nM	++++ IC50: 1.0 nM	++ IC50: 60 nM	
Nintedanib		++ IC50: 69 nM	++ IC50: 37 nM	++ IC50: 108 nM	+ IC50: 610 nM	VEGFR3, VEGFR2, LCK
PD173074		++ IC50: ~25 nM				VEGFR2,c-Src
Dovitinib		+++ IC50: 8 nM		+++ IC50: 9 nM		FLT3,c-Kit,VEGFR3/FLT4
AZD4547		++++ IC50: 0.2 nM	++++ IC50: 2.5 nM	++++ IC50: 1.8 nM		KDR
Danusertib		++ IC50: 47 nM				Aurora A, Abl, TrkA
Orantinib		+ Κι: 1.2 μΜ				PDGFRβ,Flk1
Brivanib		+ IC50: 148 nM				VEGFR2,Flk1,VEGFR1
Dovitinib Dilactic Acid		+++ IC50: 8 nM		+++ IC50: 9 nM		FLT3,c-Kit,VEGFR3/FLT4
MK-2461		+ IC50: 65 nM	++ IC50: 39 nM	+ IC50: 50 nM		c-Met (M1250T),c-Met (Y1235D), c-Met (Y1230H)
Brivanib Alaninate		+ IC50: 148 nM				VEGFR2,Flk1,VEGFR1
SSR128129E		+ IC50: 1.9 μM				
BLU-554					+++ IC50: 5 nM	
SU5402		++ IC50: 30 nM				VEGFR2,PDGFRβ
BLU9931				+ IC50: 150 nM	++++ IC50: 3 nM	
FIIN-2		+++ IC50: 3.09 nM	+++ IC50: 4.3 nM	++ IC50: 27 nM	++ IC50: 45.3 nM	
Dovitinib Lactate		+++ IC50: 8 nM		+++ IC50: 9 nM		FLT3,c-Kit,VEGFR3/FLT4
CH5183284		++ IC50: 9.3 nM	+++ IC50: 7.6 nM	++ IC50: 22 nM	+ IC50: 290 nM	
LY2874455		++++ IC50: 2.8 nM	++++ IC50: 2.6 nM	+++ IC50: 6.4 nM	+++ IC50: 6 nM	VEGFR2
Erdafitinib	V					

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com

Notes:

S2183 BGJ398 (NVP-BGJ398)

BGJ398 (NVP-BGJ398) is a potent and selective FGFR inhibitor for FGFR1/2/3 with IC50 of 0.9 nM/1.4 nM/1 nM in cell-free assays, >40-fold selective for FGFR versus FGFR4 and VEGFR2, and little activity to Abl, Fyn, Kit, Lck, Lyn and Yes. Phase 2.

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

S2801 AZD4547

Size

AZD4547 is a novel selective FGFR inhibitor targeting FGFR1/2/3 with IC50 of 0.2 nM/2.5 nM/1.8 nM in cell-free assays, weaker activity against FGFR4, VEGFR2(KDR), and little activity observed against IGFR, CDK2, and p38. Phase 2/3. 5 mg 50 mg 100 mg 10 mM/1 mL

ppias

S7665 CH5183284 (Debio-1347)

CH5183284 is a selective and orally available FGFR inhibitor with $IC_{\mbox{\tiny 50}}$ of 9.3 nM, 7.6 nM, 22 nM, and 290 nM for FGFR1, FGFR2, FGFR3, and FGFR4, respectively. Phase 1.

Size 5 mg 25 mg 100 mg

Jigo.

Other

VEGFR2/KDR,c-Met,VEGFR3/FLT4

VEGER1 VEGER2 VEGER3

FLT3,FGFR1,VEGFR3/FLT4

I vn B PDGERg PDGERß

VEGFR2.VEGFR3.EphB2

FLT1 KDR CSE-1R

PDGFRa (V561D), FLT3 (D835Y)

VEGFR1, VEGFR2, VEGFR2/Flk1

VEGFR2,PDGFRa,FGFR2

VEGFR3, VEGFR2, PDGFRa

VEGER1 VEGER2 VEGER3

FLT3,FGFR1,VEGFR3/FLT4

PDGFR,FGFR (Swiss 3T3)

FLT3 EGER1 VEGER3/ELT4

PDGFRB,Flt3,VEGFR-2

FLT3,PDGFRβ,VEGFR2

FLT3 ,PDGFRβ ,VEGFR2

S7667 SU5402

SU540 IC₅₀ of β, resp	2 is a p 20 nM, pectively	otent multi-targeted receptor tyrosine kinase 30 nM, and 510 nM for VEGFR2, FGFR1,	e inhibitor with and PDGF-R
Size	10 mg	50 mg	The second secon

S8503 BLU-554 (BLU554)

BLU-554 is a potent, highly-selective, oral FGFR4 inhibitor with an IC50 value of 5 nM. The IC50s for FGFR1-3 is 624-2203 nM.

c-Kit

IC50: 100 nM

IC50: 4.6 nM

IC50: 140 nM

IC50: 730 nM

ICco: 200 nM

IC50: 78 nM

++ IC50: 10 nM

+++

IC50: 8 nM

ICso: 80 nM

IC50: 40 nM

IC.:..: 140 nM

IC50: 1.8 µM

IC50: 37-79 nM Abl ,Src

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

concentrations of each inhibitor, please visit the website of www.selleckchem.com.

+++ IC50: 1 nM

+++ IC50: 2 nM

++++ ICso: 2 nM

+++ IC50: 9 nM

++++ ICso: 1.7 nM

+++ IC50: 2 nM

ICso: 37-79 nM Abl Src

PDGFR.v-Abl

Size 2 mg 5 mg 25 mg

c-Kit Inhibitors

Inhibitory Selectivity

Inhibitor Name

Imatinib Mesylate

Cabozantinib

Pazopanib HCI

Vatalanib 2HCI

Axitinib

Dovitinib

Masitinih

Tivozanib

Amuvatinib

OSI-930

Ki8751

Telatinib

Pazopanib

Dovitinib Dilactic Acid

Tyrphostin AG 1296

Dovitinib Lactate

Sunitinib Malate

AZD2932

Sunitinih

Notes:

specific value.

Dasatinib Monohydrate

Motesanib Diphosphate

Dasatinih

S1220 OSI-930

S1064 Masitinib (AB1010)

9737-461

Product Citations (4): Mol Syst Biol, 2015, 11(1): 789 Biomaterials, 2013, 34(38): 9737-46

Data from [Biomaterials, 2013, 34(38);

Masitinib is a novel inhibitor for Kit and PDGFRa/ß with IC50 of 200 nM

and 540 nM/800 nM, weak inhibition to ABL and c-Fms. Phase 3.

10 mg 25 mg 200 mg 10 mM/1 mL

S1244 Amuvatinib (MP-470, HPK 56)

Amuvatinib (MP-470) is a potent and multi-target inhibitor of c-Kit, PDGFRα and Flt3 with IC50 of 10 nM, 40 nM and 81 nM, respectively. Phase 2.

Product Citations (4): Nat Genet, 2012, 44(8): 852-60 Cancer Res, 2014, 74(20):		
Nat Genet, 2012, 44(8): 852-60 Cancer Res, 2014, 74(20):	Product Citation	s (4):
Cancer Res, 2014, 74(20):	Nat Genet, 2012,	44(8): 852-60
	Cancer Res, 201	4, 74(20):

Data from [Nat Genet, 2012, 44 (8):

852-601 MP-470 purchased from Selleck

ALK Inhibitors

Inhibitory Selectivity Inh

Inhibitor Name	ALK	Other
Crizotinib	+ ICso: 24 nM	c-Met
TAE684	++ IC ₅₀ : 3 nM	
Alectinib	++ IC50: 1-3.5 nM	
Ceritinib	++++ IC50: 0.2 nM	Insulin Receptor,IGF-1R
AP26113-analog	++++ IC50: 0.07 nM	EGFR(C797S/del19),IGF1R,EGFR(del19)
GSK1838705A	+++ IC ₅₀ : 0.5 nM	Insulin Receptor, IGF-1R

www.selleckchem.com

PD173074 is a potent FGFR1 inhibitor with IC50 of ~25 nM and also

inhibits VEGFR2 with IC50 of 100-200 nM in cell-free assays, ~1000-fold

Product Citations (17):

Science, 2011, 331(6019): 912-6

Cancer Discov, 2013, 3(6): 636-47

FGFR1 selective

S1264 PD173074

Size

100

80

Size

Excellent Validation, Technical Support and Prompt Delivery

selective for FGFR1 than PDGFR and c-Src.

K1

5 mg 10 mg 50 mg 10 mM/1 mL

BLU9931 is a potent, selective, and irreversible FGFR4 inhibitor with IC50 of 3 nM, about 297-, 184-, and 50-fold selectivity over FGFR1/2/3, respectively Size 5 mg 25 mg

Inhibitory Selectivity

Inhibitor Name	ALK	Other
AZD3463	+++ K:: 0.75 nM	
ASP3026	+ ICso: 3.5 nM	
Brigatinib	+++ IC50: 0.37~1.9 nM	FLT3,IGF1R,EGFR(C797S/del19)
Lorlatinib	++++ K _i : <0.07 nM	LTK (TYK1),FER,FES (FPS)
Entrectinib	1	TrkC,TrkB,TrkA

Notes:

1. For more details, such as half maximal inhibitory concentrations (IC50S) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com.

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without

specific value

S1108 TAE684 (NVP-TAE684)

TAE684 (NVP-TAE684) is a potent and selective ALK inhibitor with IC50 of 3 nM in a cell-free assay, 100-fold more sensitive for ALK than InsR. Size 5 mg 10 mg 50 mg

S7083 Ceritinib (LDK378)

Ceritinib (LDK378) is a potent inhibitor against ALK with IC50 of 0.2 nM in cell-free assays, showing 40- and 35-fold selectivity against IGF-1R and InsR, respectively. Phase 3.

CH5424802 purchased from Selleck

Size 5 mg 50 mg

S7000 AP26113

AP26113 is a potent ALK inhibitor with IC50 of 0.62 nM in a cell-free assay, demonstrated ability to overcome Crizotinib resistance mediated by a L1196M mutation. Phase 2.

Size 5 mg 10 mg 10 mM/1 mL

S7536 PF-06463922

PF-06463922 is a potent, dual ALK/ROS1 inhibitor with Ki of <0.02 nM, <0.07 nM, and 0.7 nM for ROS1, ALK (WT), and ALK (L1196M), respectively. Phase 1.

Size 5 mg 25 mg

Trk Receptor Inhibitors

S7519 GNF-5837

GNF-5837 is a selective, and orally bioavailable pan-TRK inhibitor for TrkA, and TrkB with IC50 of 8 nM, and 12 nM, respectively. Size 10 mg 50 mg 200 mg

S7998 Entrectinib (RXDX-101)

Entrectinib (RXDX-101) is an orally bioavailable pan-TrkA/B/C, ROS1 and ALK inhibitor with IC50 ranging between 0.1 and 1.7 nM. Phase 2. Size 5 mg 25 mg 100 mg

S7960 Larotrectinib (LOXO-101) sulfate new

Larotrectinib (LOXO-101) sulfate is an oral potent and selective ATP-competitive inhibitor of tropomyosin receptor kinases (TRK). Size

Ephrin Receptor Inhibitor

S2202 NVP-BHG712

Size

서신

NVP-BHG712 is a specific EphB4 inhibitor with ED50 of 25 nM that discriminates between VEGFR and EphB4 inhibition; also shows activity against c-Raf, c-Src and c-Abl with IC50 of 0.395 µM, 1.266 µM Alectinib (CH5424802) is a potent ALK inhibitor with IC50 of 1.9 nM in and 1.667 µM, respectively. cell-free assays, sensitive to L1196M mutation and higher selectivity for

5 mg 10 mg 50 mg 10 mM/1 mL Product Citation (1): NVP-BHG712 0.29uM Anticancer Research, 2014, 34(6) ▲ PD-173074 >1uM 2913-8 x I I Data independently produced by one rustome [cmpd] N NVP-BHG712 purchased from Selleck

Excellent Validation, Technical Support and Prompt Delivery

CSF-1R Inhibitors

Inhibitory Selectivity

lining	bereeting	
Inhibitor Name	CSF-1R	Other
Linifanib	+++ IC50: 3 nM	VEGFR1/FLT1,FLT3,VEGFR2/KDR
OSI-930	++ IC ₅₀ : 15 nM	FLT1,KDR,LCK
GW2580	+ IC ₅₀ : 30 nM	
CEP-32496	+++ Kd: 9 nM	c-Kit,RET,PDGFRβ
Pexidartinib	++ IC50: 20 nM	Kit,Flt3
BLZ945	++++ IC50: 1 nM	
Notes:		

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

S8042 GW2580 (SC-203877)

GW2580 is a selective CSF-1R inhibitor for c-FMS with IC50 of 30 nM, 150- to 500-fold selective compared to b-Raf, CDK4, c-KIT, c-SRC, EGFR, ERBB2/4, ERK2, FLT-3, GSK3, ITK, JAK2 etc. , State, Size 25 mg 10 mM/1 mL

S1003 Linifanib (ABT-869, AL39324, RG3635)

Linifanib (ABT-869) is a novel, potent ATP-competitive VEGFR/PDGFR inhibitor for KDR, CSF-1R, FIt-1/3 and PDGFRβ with IC50 of 4 nM, 3 nM, 3 nM/4 nM and 66 nM respectively, mostly effective in mutant kinase-dependent cancer cells (i.e. FLT3). Phase 3. Page 37

S7725 BLZ945

BLZ945 is an orally active, potent and selective CSF-1R inhibitor with IC50 of 1 nM, >1000-fold selective against its closest receptor tyrosine kinase homologs.

Size 5 mg 25 mg 100 mg

to at

S7818 Pexidartinib (PLX3397)

Pexidartinib (PLX3397) is an oral, potent mutil-target receptor tyrosine kinase inhibitor of CSF-1R. Kit. and Flt3 with IC50 of 20 nM. 10 nM and 160 nM, respectively. Phase 3. and and the

10 mg 50 mg Size

TAM Receptor Inhibitors

Inhibitory Selectivity

Inhibitor Name	Mer	Axi	Tyro3	Other
BMS-777607	++ IC50: 14 nM	++++ IC50: 1.1 nM	++++ IC50: 4.3 nM	RON,Met,FLT3
R428		++ IC50: 14 nM		
UNC2250	++++ IC50: 1.7 nM		+ IC ₅₀ : 100 nM	
Sitravatinib	++++ IC50: 2 nM	++++ IC50: 1.5 nM		DDR2,EPHA3,VEGFR3 (FLT4)
RXDX-106	+ IC50: 29 nM	+++ ICso: 7 nM	++ IC50: 19~29 nM	c-Met,VEGFR2
UNC2025	++++ IC ₅₀ : 0.74 nM		++ IC ₅₀ : 17 nM	FLT3
TP-0903		++ IC50: 27 nM		
NPS-1034		+++ ICso: 10.3 nM		Met
LDC1267	+++ IC50: <5 nM	+ ICso: 29 nM	+++ IC50: 8 nM	
UNC2881	+++ IC ₅₀ : 4.3 nM	+ IC ₅₀ : 360 nM	+ IC ₅₀ : 250 nM	

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsss) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

S2841 R428 (BGB324)

R428 (BGB324) is an inhibitor of Axl with IC50 of 14 nM, >100-fold selective for Axl versus Abl. Selectivity for Axl is also greater than Mer and Tyro3 (50-to-100- fold more selective) and InsR, EGFR, HER2, and PDGFRB (100- fold more selective).

S1119 Cabozantinib (XL184, BMS-907351)

Cabozantinib (XL184, BMS-907351) is a potent VEGFR2 inhibitor with IC50 of 0.035 nM and also inhibits c-Met, Ret, Kit, FIt-1/3/4, Tie2, and AXL with IC50 of 1.3 nM, 4 nM, 4.6 nM, 12 nM/11.3 nM/6 nM, 14.3 nM and 7 nM in cell-free assays, respectively.

S1561 BMS-777607

BMS-777607 is a Met-related inhibitor for c-Met. Axl. Ron and Tvro3 with IC50 of 3.9 nM, 1.1 nM, 1.8 nM and 4.3 nM in cell-free assays. 40-fold more selective for Met-related targets versus Lck. VEGFR-2. and TrkA/B, and more than 500-fold greater selectivity versus all other receptor and non-receptor kinases. Phase 1/2.

M114.038547] R428 purchased from Selleck

- 2803-11

Product Citations (5):

Cancer Res, 2014, 74(18): 5152-64 Mol Cell Proteomics, 2014, 13(11): Data from [Mol Cell Proteomics, 2014,

CSF-1R / TAM Receptor

Angiogenesis

JAK Inhibitors

EGFR Inhibitors

Detailed product information is on pa

Detailed product information is on pa

PDGFR Inhibitors

Angiogenesis

	FGFR Inhibitors	
ge 24-26	Detailed product information is on page 46-47	
	ALK Inhibitors	
ge 38-40	Detailed product information is on page 47-48	
	HIF Inhibitors	

Detailed product information is on page 27

FLT3 Inhibitors

Detailed product information is on page 45

50

Detailed product information is on page 43-44

Detailed product information is on page 41-42

V	D	A	

S1537 DMXAA (Vadimezan)

Bcr-Abl Inhibitors

Bcr-Abl

+++IC50: 0.6 nM

+++IC50: 0.37 nM

++ IC50: <30 nM

IC50: 25 nM

IC50: 1.8 µM

++ IC50: 5.8 nM

+ ICso: 14 nM

+++IC50: 0.34 nM

IC50: 273 nM

+++ IC50: 122 nM

++ ICso: 34 nM

+++IC50: 0.6 nM

+++ IC50: 1-2 nM

S1026 Imatinib Mesylate (STI571)

IC50: 220 nM

Inhibitory Selectivity

Inhibitor Name

Dasatinib

Saracatinib

Ponatinib

Nilotinib

Danusertib

AT9283

Degrasyn

Bafetinih

KW-2449

Rehastinih

G7D824

GNF-2

GNF-7

Radotinib

Dasatinib Monohydrate

GNF-5

PD173955

40

10

-c-Kit^{Y71}

-actin

Notes:

Dimesylate

NVP-BHG712

Imatinib Mesylate

DMXAA (Vadimezan) is a vascular disrupting agents (VDA) and competitive inhibitor of DT-diaphorase with K of 20 µM and IC50 of 62.5 µM in cell-free assays, respectively. Phase 3. Size 5 mg 25 mg 100 mg 10 mM/1 mL , QQQ,

Abl

++ IC50: 0.6 nM

IC50: 30 nM

++ IC50: 0.37 nM

IC50: 25 nM

++ ICso: 4-30 nM

++ IC50: 5.8 nM

++ ICso: 4-14 nM

IC50: 1.667 µM

+ IC50: 0.75-5 nM

+++ ICso: 0.75-5 nM

ICso: 133 nM

++ IC50: 0.6 nM

IC50: 600 nM

Other

Src,c-Kit (D816V),c-Kit (wt)

c-Src,LCK,EGFR (L861Q)

PDGFRα,VEGFR2,FGFR1

Aurora A.TrkA.RET

JAK3.JAK2.Aurora B

FLT3 (D835Y),FLT3,FGFR1

Src,c-Kit (D816V),c-Kit (wt)

EphB4,C-Raf,c-Src

FLT3,KDR,Tie-2

DUB

Lyn

-Kit PDGFR

S1490 Ponatinib (AP24534)

Ponatinib (AP24534) is a novel, potent multi-target inhibitor of Abl, PDGFRα, VEGFR2, FGFR1 and Src with IC50 of 0.37 nM, 1.1 nM, 1.5 nM, 2.2 nM and 5.4 nM in cell-free assays, respectively.

Size 10 mg 50 mg 200 mg 10 mM/1 mL

S1033 Nilotinib (AMN-107)

Size

Nilotinib (AMN-107) is a selective Bcr-Abl inhibitor with IC₅₀ less than 30 nM in Murine myeloid progenitor cells.

3661-731 Nilotinib (µM) 0 1 2.5 Nilotinib purchased from Selleck

S2243 Degrasyn (WP1130)

Degrasyn (WP1130) is a selective deubiquitinase(DUB: USP5, UCH-L1, USP9x, USP14, and UCH37) inhibitor and also suppresses Bcr/Abl; also a JAK2 transducer (without affecting 20S proteasome) and activator of transcription (STAT).

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Bafetinib (INNO-406) is a potent and selective dual Bcr-Abl/Lyn inhibitor with IC50 of 5.8 nM/19 nM in cell-free assays, does not inhibit the phosphorylation of the T315I mutant and is less potent to PDGFR and c-Kit. Phase 2. Mr. Hote

5 mg 25 mg 100 mg Size

S1369 Bafetinib (INNO-406, NS-187)

Product Citations (2): J Med Chem, 2015,58(1): 466-79 Int Arch Alleray Immunol, 2012, 159(1) 15-22

Data from [Int Arch Allergy Immunol, 2012, 159(1): 15-22] INNO-406 purchased from Sellect

Imatinib Mesylate (STI571) is an orally bioavailability mesylate salt of Imatinib, which is a multi-target inhibitor of v-Abl, c-Kit and PDGFR with IC50 of 0.6 µM, 0.1 µM and 0.1 µM in cell-free or cell-based assays, respectively Size 100 mg 250 mg 10 mM/1 mL

1 For more details, such as half maximal inhibitory concentrations (ICsps) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

- MelMS 2100 -0-M230 -D- WM3211 Product Citations (28): Cancer Cell, 2014, 26(6): 840-50 Cell Stem Cell, 2012, 10(2); 210-7 1114 100 1000 10000 Imatinib mesylate (nM) otal c-Ki

Data from [Oncogene, 2012, 33(2): 236-451 Imatinib Mesylate purchased from Selleck

S1134 AT9283

AT9283 is a potent JAK2/3 inhibitor with IC50 of 1.2 nM/1.1 nM in cell-free assays; also potent to Aurora A/B, Abl(T315I). Phase 2.

Page 25

S2158 KW-2449

KW-2449 is a multiple-target inhibitor, mostly for FIt3 with IC $_{50}$ of 6.6 nM, modestly potent to FGFR1, Bcr-Abl and Aurora A; little effect on PDGFR β, IGF-1R, EGFR. Phase 1. ·---- Page 45

Src Inhibitors

Inhibitory Selectivity

Inhibitor Name	Src	Lck	Fyn	Lyn	Yes	Other
Dasatinib	++++ IC50: 0.8 nM					Abl,c-Kit (D816V),c-Kit (wt)
Saracatinib	++++ IC50: 2.7 nM	++++ IC ₅₀ : <4 nM	++ IC50: 10 nM	+++ IC50: 5 nM		EGFR (L861Q),c-YES,EGFR (L858R)
Bosutinib	++++ IC50: 1.2 nM					Abl
KX2-391	++ GI50: 9~60 nM					
NVP-BHG712	+ IC ₅₀ : 1.266 μM					EphB4,C-Raf,c-Abl
PP2	+++ IC50: 5 nM	+++ IC50: 4 nM	+++ IC ₅₀ : 5 nM			EGFR
PP1	+++ IC50: 6 nM	+++ IC50: 5 nM	++ IC50: 6 nM			Kit,EGFR,Bcr-Abl
SU6656	+ IC50: 280 nM		+ ICso: 170 nM	+ ICso: 130 nM	++ IC50: 20 nM	
Dasatinib Monohydrate	++++ IC50: 0.8 nM					Abl ,c-Kit (D816V),c-Kit (wt)
WH-4-023	++++ IC50: 6 nM	++++ IC50: 2 nM				
Quercetin	1					Sirtuin,PKC,PI3Ky

S1107 Danusertib (PHA-739358)

CDK2 etc. Phase 2.

Danusertib (PHA-739358) is an Aurora kinase inhibitor for Aurora A/B/C

with IC50 of 13 nM/79 nM/61 nM in cell-free assays, modestly potent to

Abl, TrkA, c-RET and FGFR1, and less potent to Lck, VEGFR2/3, c-Kit,

Page 28

Notes:

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Angioger

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

S7565 WH-4-023

WH-4-023 is a potent and orally active Lck/Src inhibitor with IC50s of 2 nM and 6 nM in cell-free assays, respectively. Exhibits >300-fold selectivity against p38α and KDR. Also potently inhibits SIK (IC₅₀ values are 10, 22 and 60 nM for SIK 1, 2 and 3 respectively) and displays selectivity over a range of closely related kinases.

Size 5 mg 25 mg 100 mg

S7060 PP1

PP1 is a potent and selective Src inhibitor for Lck/Fyn with IC50 of 5 nM/ 6 nM.

Size 10 mg 25 mg

S2391 Quercetin (Sophoretin)

Quercetin, a natural flavonoid present in vegetables, fruit and wine, is a stimulator of recombinant SIRT1 and also a PI3K inhibitor with IC50 of 2.4-5.4 uM. Phase 4. ----- Page 30

S7774 SU6656

SU6656 is a selective Src family kinase inhibitor with IC50 of 280 nM, 20 nM, 130 nM, and 170 nM for Src, Yes, Lyn, and Fyn, respectively.

Size 5 mg 25 mg 100 mg

S7782 Dasatinib Monohydrate

Syk Inhibitors

Inhibitory Selectivity

Svk

++ IC50; 41 nM

++ IC50: 41 nM

++ IC50: 41 nM

+++ IC50: 1 nM

++ ICso: 41 nM

++++ IC50: 4nM

+++ IC50: 7.7 nM

+++ IC50: 5.6 nM

+++ K_i: 7.5 nM

ICso: 2.5 µM p97.Src

concentrations of each inhibitor, please visit the website of www.selleckchem.com.

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

3. Red "V" refers to compounds which do inhibitory effects on the related isoform, but without

Dasatinib Monohydrate is a novel, potent and multi-target inhibitor that targets Abl, Src and c-Kit, with IC50 of <1 nM, 0.8 nM and 79 nM, respectively.

Size 50 mg 200 mg

Inhibitor Name

R788 Disodium

PRT062607 HCI

Fostamatinib

PRT-060318

Entospletinib

BAY-61-3606

Piceatannol

specific value.

Notes:

RO9021

R406

R406

MNS

S8032 PRT062607 (P505-15, BIIB057) HCI

S7523 Entospletinib (GS-9973)

PRT062607 (P505-15) HCl is a novel, highly selective Syk inhibitor with IC50 of 1 nM in cell-free assays, >80-fold selective for Syk than for Fgr, Lyn, FAK, Pyk2 and Zap70. Size 5 mg 25 mg 10 mM/1 mL

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Size 10 mg 50 mg 200 mg

response.

S3026 Piceatannol

Piceatannol, a natural stilbene, is a selective Syk inhibitor and ~10-fold selectivity versus Lyn.

Size 10 mg 25 mg 50 mg 10 mM/1 mL

S7286 RO9021

RO9021 potently inhibits SYK kinase activity with an average IC50 of 5.6 nM and suppresses B-cell receptor signaling.

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working Size 1 mg 5 mg 25 mg

strongly inhibiting Syk but not Lyn, 5-fold less potent to Flt3. Phase 1.

Src / Syk

waat 😽 Product Citations (15): Immunity, 2014, 40(3); 389-99 Nat Cell Biol. 2015. 17(1): 57-67 Data from [Blood, 2013, 122(4); 580-91 R406 purchased from Selleck

R406 is a potent Syk inhibitor with IC50 of 41 nM in cell-free assays,

2 mg 10 mg 50 mg 10 mM/1 mL

S2206 R788 (Fostamatinib) Disodium

R788 (Fostamatinib) disodium, a prodrug of the active metabolite R406, is a Syk inhibitor with IC50 of 41 nM in cell-free assays, strongly inhibits

S2194 R406

Size

N

Ϋ́Ο

ni CCCP

N-N-N-

Other

Adenosine A3 receptor, Adenosine transporte

FGR.MLK1.YES

Lyn,PKA,PKC

Monoamine transporter

 $\propto_{\tilde{D}}$

R406 (free base) is a potent Syk inhibitor with IC50 of 41 nM in a cell-free assay, strongly inhibits Syk but not Lyn, 5-fold less potent to Flt3. Phase

FAK / BTK

FAK Inhibitors

Inhibitory Selectivity

Inhibitor Name	FAK	Other
PF-00562271	++++ IC50: 1.5 nM	CDK2/CyclinE,CDK3/CyclinE,CDK1/CyclinB
PF-562271	++++ IC50: 1.5 nM	CDK2/CyclinE,CDK3/CyclinE,CDK1/CyclinB
PF-573228	+ IC50: 4 nM	
TAE226	++ IC50: 3.5~5.5 nM	Insulin Receptor,IGF-1R,c-Met
PF-03814735	+ IC50: 22 nM	Aurora A, Aurora B, FLT1
PF-562271 HCI	++++ IC50: 1.5~13 nM	CDK2/CyclinE,CDK3/CyclinE,CDK1/CyclinB
GSK2256098	++++ K: 0.4 nM	
PF-431396	++ IC50: 2~11 nM	
PND-1186	++++ IC50: 0.5 nM	
Defactinib	V	

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

S2672 PF-00562271

PF-00562271 is the benzenesulfonate salt of PF-562271, which is a potent, ATP-competitive, reversible inhibitor of FAK with IC50 of 1.5 nM, ~10-fold less potent for Pyk2 than for FAK and >100-fold selectivity against other protein kinases, except for some CDKs. Phase 1.

Size 5 mg 10 mg D FAK inh 125 100 Product Citation (1): Mol Ther. 2012. 20(5): 972-83 Data from [Mol Ther, 2012, 20(5): 972-83] PF-00562271 (FAK inhibitor) purchased Mn++ from Selleck

S2890 PE-562271

PF-562271 is a potent, ATP-competitive, reversible inhibitor of FAK with IC50 of 1.5 nM in cell-free assays, ~10-fold less potent for Pyk2 than for FAK and >100-fold selectivity against other protein kinases, except for some CDKs

	OVIOL	
\$2013	DF-573228	
02013	11-5/5220	

PF-573228 is an ATP-competitive inhibitor of FAK with IC50 of 4 nM in a cell-free assay, ~50- to 250-fold selective for FAK than for Pyk2, CDK1/7 and GSK-3β.

e885871

PF-562271 purchased from Selleck

Product Citations (4): J Cell Sci, 2014, 127(Pt 14): 3039-51

Data from [PLoS One, 2014, 9(2);

PF-573228 purchased from Selleck

J Biol Chem, 2015,

e885871

10 1074/ibc M114 624247

Size 10 mg 50 mg 10 mM/1 mL

S7653 PND-1186 (VS-4718) PND-1186 (VS-4718) is a reversible and selective FAK inhibitor with IC50 of 1.5 nM. Phase 1. Size 5 mg 25 mg 100 mg S7654 Defactinib (VS-6063, PF-04554878) Defactinib (VS-6063, PF-04554878) is a selective, and orally active FAK inhibitor. Phase 2. Size 5 mg 25 mg 100 mg S2725 PF-03814735 PF-03814735 is a novel, potent and reversible inhibitor of Aurora A/B with IC50 of 0.8 nM/5 nM, is less potent to FIt3, FAK, TrkA, and minimally active to Met and FGFR1. Phase 1. ----- Page 29 C^YM S8523 GSK2256098 GSK2256098 is a potent, selective, reversible, and ATP competitive FAK kinase inhibitor with apparent K of 0.4 nM. Size 5 mg 25 mg

S2820 TAE226 (NVP-TAE226)

5 mg 10 mg 10 mM/1 mL

IGF-1R, ALK, and c-Met.

Size

TAE226 (NVP-TAE226) is a potent FAK inhibitor with IC50 of 5.5 nM and

modestly potent to Pyk2, ~10- to 100-fold less potent against InsR,

LBH589

Product Citations (2):

new

e1134]

Cell Death Dis. 2014. 5: e1134 J Biol Chem, 2015, 290(14): 8677-92

Data from [Cell Death Dis. 2014. 5:

TAE226 purchased from Selleck

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Notes:

Excellent Validation, Technical Support and Prompt Delivery

BTK Inhibitors Inhibitory Selectivity Inhibitor Name Othor

Infilbitor Name		DIK	Other
Ibrutinib	++++	IC50: 0.5 nM	BLK,Bmx,FGR
AVL-292	++++	IC ₅₀ : <0.5 nM	
CNX-774	+++	IC ₅₀ : <1 nM	
Acalabrutinib	++	IC50: 3 nM	
ONO-4059 analogue	++	IC50: 23.9 nM	
LFM-A13	+	K _i : 1.4 μM	
RN486	++	IC50: 4 nM	
CGI1746	+++	IC50: 1.9 nM	

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

S2680 Ibrutinib (PCI-32765)

Ibrutinib (PCI-32765) is a potent and highly selective Brutons tyrosine kinase (Btk) inhibitor with IC50 of 0.5 nM in cell-free assays, modestly potent to Bmx, CSK, FGR, BRK, HCK, less potent to EGFR, Yes, ErbB2, JAK3 etc.

S7173 CC-292 (AVL-292)

CC-292 (AVL-292) is a covalent, orally active, and highly selective BTK inhibitor with IC50 of <0.5 nM, displaying at least 1400-fold selectivity over the other kinases assayed. Phase 1.

Acalabrutinib(ACP-196) is a selective second-generation Bruton's tyrosine kinase (BTK) inhibitor, which prevents the activation of the B-cell antigen receptor (BCR) signaling pathway. ACP-196 has improved target specificity over ibrutinib with 323-, 94-, 19- and 9-fold selectivity over the other TEC kinase family members (ITK, TXK, BMX, and TEC, respectively) and no activity against EGFR.

Size 5 mg 25 mg 100 mg

55

Apoptosis

Bcl-2 Inhibitors Activator

Inhibitory Selectivity

Inhibitor Name	Bcl-2	BcI-B	Bcl-w	Bcl-xL	McI-1	Bax	Other
ABT-737	+++ EC50: 30.3 nM	+ EC50: 1.82 μM	+++ EC50: 197.8 nM	+++ EC50: 78.7 nM			
Navitoclax (ABT-263)	++++ K _i : ≤1 nM		++++ K _i : ≤1 nM	++++ Ki: ≤0.5 nM	++ K _i : 550 nM		
Obatoclax Mesylate	+++ K _i : 0.22 μΜ						
TW-37	++ Ki: 0.29 μM			+ Κ _i : 1.11 μΜ	+++ Ki: 0.26 μM		
Venetoclax	++++Ki: <0.01 nM						
AT101	++ K _i : 0.32 μM			++ Ki: 0.48 μM	+++ K _i : 0.18 μM		
HA14-1	+ IC ₅₀ : 9 μΜ						
Sabutoclax	++ IC ₅₀ : 0.32~0.62 μM			++ IC ₅₀ : 0.31 μM	+++ IC ₅₀ : 0.20 μM	++ IC50: 0.62 μM	
A-1155463				++++ Ki: <0.01 nM			
A-1210477					++++ IC ₅₀ : 26.2 nM		
UMI-77					++ Ki: 490 nM		
Gambogic Acid	+ IC50: 1.06~1.21 μM	++ IC50: 0.66 μM	++++ IC50: 0.02 μM	+ IC ₅₀ : 1.47 μM	+ ICso: 0.79 μM	+ IC50: 1.06 μM	Caspase

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

Bcl-2 selective

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Bcl-2 Inhibitors

S1002 ABT-737

ABT-737 is a BH3 mimetic inhibitor of Bcl-xL. Bcl-2 and Bcl-w with EC50 of 78.7 nM, 30.3 nM and 197.8 nM in cell-free assays, respectively; no inhibition observed against Mcl-1, Bcl-B or Bfl-1. Phase 2. 5 mg 50 mg 100 mg 10 mM/1 mL Size

S1001 Navitoclax (ABT-263)

Navitoclax (ABT-263) is a potent inhibitor of Bcl-xL. Bcl-2 and Bcl-w with K of \leq 0.5 nM, \leq 1 nM and \leq 1 nM in cell-free assays, but binds more weakly to Mcl-1 and A1. Phase 2.

Data from [Cancer Res, 2012, 72 (12): 2949-561 ABT-263 purchased from Selleck

S1057 Obatoclax Mesylate (GX15-070)

Obatoclax Mesylate (GX15-070) is an antagonist of Bcl-2 with K of 0.22 µM in a cell-free assay, can assist in overcoming MCL-1 mediated resistance to apoptosis. Phase 3. сн,воун

Size 5 mg 10 mg 50 mg 10 mM/1 mL

S1121 TW-37

TW-37 is a novel non-peptide inhibitor to recombinant Bcl-2, Bcl-xL and Mcl-1 with K of 0.29 $\mu\text{M},$ 1.11 μM and 0.26 μM in cell-free assays, respectively. the sport

Sabutoclax (BI-97C1) is a pan-Bcl-2 inhibitor, including Bcl-xL, Bcl-2,

Mcl-1 and Bfl-1 with IC50 of 0.31 µM, 0.32 µM, 0.20 µM and 0.62 µM,

Size 10 mg 50 mg 10 mM/1 mL

S8061 Sabutoclax (BI-97C1)

respectively. Size 5 mg 50 mg 20(11): 1475-841

Product Citations (12):

Nat Chem Biol, 2015, 10.1038/nchembio.1797

Data from [Cell Death Differ, 2013.

Cell Death Differ, 2013, 20(11); 1475-84

TW-37 purchased from Selleck

www.selleckchem.com

Inhibitory Selectivity P 00 Inhibitor Name Caspase Caspase-1 Caspase-3 Caspase-4 Belnacasa ++++ Ki: 0.8 nN ++++ K_i: <0.6 nM Emricasar 1 7-VAD-EMK Z-DEVD-FMK

Notes: 1. For more details, such as half maximal inhibitory concentrations (IC50s) and working

concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

Caspase Inhibitors Activator

Cancer Res, 2011, 71(13): 4494-505 Data from [Cell Death Dis. 2012. 3: e351]

GX15-070 purchased from Selleck

A-1155463, a highly potent and selective BCL-XL inhibitor, shows picomolar binding affinity to BCL-XL, and >1000-fold weaker binding to

Bcl-2 Activator

Size 10 mg 50 mg

S8048 Venetoclax (ABT-199, GDC-0199)

5 mg 50 mg 10 mM/1 mL

Size

CLL only

S7531 UMI-77

5 mg 25 mg

S7790 A-1210477

Bcl-2 family members.

Size 5 mg 25 mg

S7800 A-1155463

Size

Size

BcI-2 selective

CQ-CL

and Bcl-w, and no activity to Mcl-1. Phase 3.

CLL on stroma

selectivity over other members of Bcl-2 family.

Venetoclax (ABT-199, GDC-0199) is a Bcl-2-selective inhibitor with Ki of <0.01 nM in cell-free assays, >4800-fold more selective versus Bcl-xL

> Product Citations (11): Leukemia, 2014, 28(8); 1657-65

Cell Death Differ. 2015.

Data from [J Biol Chem, 2014, 289(23):

ABT-199 purchased from Selleck

10.1038/cdd.2015.73

16190-91

UMI-77 is a selective McI-1 inhibitor with K of 490 nM, showing

A-1210477 is a potent and selective MCL-1 inhibitor with Ki and IC50 of 0.454 nM and 26.2 nM, respectively, >100-fold selectivity over other

S7105 BAM7 BAM7 is a direct and selective activator of pro-apoptotic Bax with EC50 of 3.3 µM.

Caspase Inhibitors

S7023 Z-VAD-FMK

Z-VAD-FMK is a cell-permeable, irreversible pan-caspase inhibitor. blocking all features of apoptosis in THP.1 and Jurkat T-cells. Size 1 mg 5 mg

S2228 Belnacasan (VX-765)

Belnacasan (VX-765) is a potent and selective inhibitor of caspase-1 with K of 0.8 nM in a cell-free assay. Phase 2. Size 10 mg 50 mg PULLO"

Caspase-1 selective

S7312 Z-DEVD-FMK

Z-DEVD-FMK is a specific, irreversible Caspase-3 inhibitor, and also shows potent inhibition on caspase-6, caspase-7, caspase-8, and caspase-10. Size 1 mg an hall the

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S7775 Emricasan
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Emricasan is a potent irreversible pan-caspase inhibitor.

Size 5 mg 25 mg

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new

Caspase Activator

S2738 PAC-1

PAC-1 is a potent procaspase-3 activator with EC $_{50}$ of 0.22 μM and the first small molecule known to directly activate procaspase-3 to caspase-3.

Size 10 mg 50 mg 250 mg 10 mM/1 mL

p53 Inhibitors | Activators

Apoptosis p53 Inhibitors

S2929 Pifithrin-α (PFTα)

Pifithrin-a is an inhibitor of p53, inhibiting p53-dependent transactivation of p53-responsive genes. Size 25 mg 50 mg 10 mM/1 mL $(-)^{(-)}$

S2930 Pifithrin-u

58

Pifithrin-µ is a specific p53 inhibitor by reducing its affinity to Bcl-xL and Bcl-2, and also inhibits HSP70 function and autophagy. 0 Size 10 mg 50 mg

Excellent Validation, Technical Support and Prompt Delivery

S2781 RITA (NSC 652287)

p53 Activators

S1172 JNJ-26854165 (Serdemetan)

RITA (NSC 652287) induces both DNA-protein and DNA-DNA cross-links with no detectable DNA single-strand breaks, and also inhibits MDM2-p53 interaction by targeting p53. Size 5 mg 10 mg 10 mM/1 mL

С С С С

JNJ-26854165 (Serdemetan) acts as a HDM2 ubiguitin ligase antagonist and also induces early apoptosis in p53 wild-type cells, inhibits cellular proliferation followed by delayed apoptosis in the absence of functional p53. Phase 1. 5 mg 25mg 100 mg 10 mM/1 mL No treatment Serdemetan Product Citations (2): Sci Rep, 2014, 4: 4664 Head Neck, 2014, 10.1002/hed.23822

Data from [Sci Rep. 2014 4: 4664] erdemetan purchased from Sellecl

S7149 NSC 319726

TNF-alpha Inhibitors Inhibitory Selectivity

Inhibitor Name	TNF-α	Other
Pomalidomide	+++ IC50: 13 nM	
Necrostatin-1	+ EC ₅₀ : 490 nM	
QNZ	+++ IC50: 7 nM	NF-ĸB
Thalidomide	1	
Notes:		

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

S1567 Pomalidomide

O.L.D.

Pomalidomide inhibits LPS-induced TNF- α release with IC₅₀ of 13 nM in PBMCs.

Thalidomide was introduced as a sedative drug, immunomodulatory agent and also is investigated for treating symptoms of many cancers. Thalidomide inhibits an E3 ubiquitin ligase, which is a CRBN-DDB1-Cul4A complex. Size 200 mg ~~~~

S8037 Necrostatin-1

Necrostatin-1 is a specific RIP1 inhibitor and inhibits TNF-α-induced necroptosis with EC50 of 490 nM in 293T cells. CO (1 - s

Size 10 mg 100 mg 10 mM/1 mL

S1623 Acetylcysteine

Acetylcysteine(N-acetyl-I-cysteine) is a ROS(reactive oxygen species) inhibitor that antagonizes the activity of proteasome inhibitors. It is also a tumor necrosis factor production inhibitor, used mainly as a mucolytic, protects against acetaminophen overdose-induced hepatotoxicity by maintaining or restoring hepatic concentrations of glutathione.

Size 10 mg 50 mg 10 mM/1 mL

S4902 QNZ (EVP4593)

QNZ (EVP4593) shows potent inhibitory activity toward both NF-KB activation and TNF-α production with IC50 of 11 nM and 7 nM in Jurkat T cells, respectively,

Mdm2 Inhibitors | Activator | Antagonists

Inhibitory Selectivity

Inhibitor Name	Mdm2	Othe
Nutlin-3	++ IC50: 180 nM	
Nutlin-3a	++ IC ₅₀ : 90 nM	
Nutlin-3b	+ ICso: 13.6 μM	
MX69	+ Kd: 2.34 μM	
MI-773 (SAR405838)	++++ IC50: 0.88 nM	p53
Idasanutlin (RG-7388)	++++ IC50: 6 nM	
RG-7112	+++ Kd: 11 nM	
YH239-EE	1	

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "V" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

Mdm2 Inhibitors

S8059 Nutlin-3a

Nutlin-3a, the active enantiomer of Nutlin-3, inhibits the p53/MDM2 interaction with IC50 of 90 nM in a cell-free assav. Size 5 mg 25 mg

Idasanutlin (RG-7388) is a potent and selective p53-MDM2 inhibitor with IC₅₀ of 6 nM showing improved in vitro binding as well as cellular potency/selectivity.

5 mg 25 mg Size

Mdm2 Activator

S2678 NSC 207895

NSC 207895 suppresses MDMX with IC50 of 2.5 µM, leading to enhanced p53 stabilization/activation and DNA damage, and also regulates MDM2, an E3 ligase.

Size 5 mg 10 mg 50 mg

Mdm2 Antagonists

S1061 Nutlin-3

Nutlin-3 is a potent and selective Mdm2 (RING finger-dependent ubiguitin protein ligase for itself and p53) antagonist with IC50 of 90 nM in a cell-free assay; stabilizes p73 in p53-deficient cells.

Size	5 mg	25 mg 100	mg 10 mM	<u>/1 mL</u>
	UKF-NB-3	UKF-NB-3 ^r NUTLIN ^{10µM}	UKF-NB-3' RITA ^{10µM}	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
	non treated RITA	non treated Autor ATA	non-treated	Product Citations (8): Hepatology, 2015, 10.1002/hep.27992
p53 pp53				int J Cancer, 2014, 10.1002/ijc.29194
p21 BAX PUMA	-		-	Data from [Cell Death Dis, 2012, 3: e294]
MDM2	_		tion was been	Nutlin-3 purchased from Selleck

S7649 MI-773 (SAR405838)

MI-773 (SAR405838) is an orally available MDM2 antagonist with Ki of 0.88 nM. Phase 1. Size 5 mg 25 mg

S7030 RG-7112

RG7112 (RO5045337) is an orally bioavailable and selective p53-MDM2 inhibitor with HTRF IC₅₀ of 18 nM. Size 5 mg 25 mg

-0R

Survivin Inhibitor

YM155

S1130 YM155 (Sepantronium Bromide)

YM155 (Sepantronium Bromide) is a potent survivin suppressant by inhibiting Survivin promoter activity with IC50 of 0.54 nM in HeLa-SURP-luc and CHO-SV40-luc cells; does not significantly inhibit SV40 promoter activity, but is observed to slightly inhibit the interaction of Survivin with XIAP. Phase 2.

Size 5 mg 25 mg 100 mg 10 mM/1 mL

*] p = <0.0001 (n = 30)</p>

4

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DMSO

* 2012, 109(2): 600-5] YM155 YM155 purchased from Selleck
IAP Inhibitors | Antagonist

Inhibitory Selectivity

Inhibitor Name	cIAP	XIAP	Other
Birinapant	++++ Kd: <1 nM	++ Kd: 45 nM	
GDC-0152	+++ Ki: 17~43 nM	++ Ki: 28~112 nM	MLXBIR3SG
Embelin		+ IC ₅₀ : 4.1 μM	5-LO,mPGES-1
BV-6	1		
LCL161	1		

Notes:

1. For more details, such as half maximal inhibitory concentrations (IC50S) and wo concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

IAP Inhibitors

S7015 Birinapant

Birinapant is a SMAC mimetic antagonist, mostly to cIAP1 with Kd of <1 nM in a cell-free assay, less potent to XIAP. Phase 2.



S7009 LCL161

I CI -161 a small molecule second mitochondrial activator of caspase (SMAC) mimetic, potently binds to and inhibits multiple IAPs (i.e. XIAP, c-IAP).

GDC-0152 is a potent antagonist of XIAP-BIR3, ML-IAP-BIR3,

cIAP1-BIR3 and cIAP2-BIR3 with K of 28 nM, 14 nM, 17 nM and 43 nM

Size 5 mg 25 mg 100 mg

S7597 BV-6

BV-6 is a SMAC mimetic, dual cIAP and XIAP inhibitor.

Size 5 mg 25 mg 100 mg

IAP Antagonist

S7010 GDC-0152

cIAP2-BIR2, Phase 1,

Size 10 mg

Apoptosis



Size

S7307 GSK2606414

GSK2606414 is an orally available, potent, and selective PERK inhibitor with IC50 of 0.4 nM, displaying at least 100-fold selectivity over the other EIF2AKs assayed

Serine/threonin Kinase

CRT0066101 is a small molecule PKD family specific inhibitor which

PERK

++++ IC50: 0.4 nM

+++ ICso: 0.9 nM

++ IC50: 5 nM

specifically blocks PKD1/2 activity and does not suppress PKCa/

Inhibitor

S8366 CRT0066101

Size 5 mg 25 mg

Inhibitor Name

PKCβ/PKCε activity in multiple.

PERK Inhibitors

Inhibitory Selectivity

5°C

- Eĥ

S7033 GSK2656157

5 mg

GSK2656157 is an ATP-competitive and highly selective inhibitor of PERK with IC50 of 0.9 nM in a cell-free assay, 500-fold greater against a panel of 300 kinases Size 50 mg

ISRIB (trans-isomer S7400

ISRIB (trans-isomer), the trans-isomer of ISRIB, is a potent and selective PERK inhibitor with IC50 of 5 nM and does not have global effects on translation, transcription, or mRNA stability in non-stressed cells Size 10 mg 25 mg

anor O

S2923 Salubrina

5 mg 10 mM/1 mL

in cell-free assays, respectively; less affinity shown to cIAP1-BIR2 and Salubrinal is a selective inhibitor of eIE2a dephosphorylation and inhibits ER stress-mediated apoptosis with EC $_{50}$ of ~15 μM in a cell-free assay

Autophagy



Autophagy Inhibitors | Activators | Modulators

Autophagy Inhibitors

S1105 LY294002

LY294002 is the first synthetic molecule known to inhibit PI3Ka/ δ/β with IC50 of 0.5 µM/0.57 µM/0.97 µM in cell-free assays, respectively; more stable in solution than Wortmannin, and also blocks autophagosome formation ----- Page 7

S1150 Paclitaxel

Paclitaxel is a microtubule polymer stabilizer with IC50 of 0.1 pM in human endothelial cells. ----- Page 76

S2758 Wortmannin

Wortmannin is the first described PI3K inhibitor with IC50 of 3 nM in a cell-free assay, with little selectivity within the PI3K family. Also blocks autophagosome formation and potently inhibits DNA-PK/ATM with IC50 of 16 nM and 150 nM in cell-free assays. Page 8

S2767 3-Methyladenine (3-MA)

3-Methyladenine (3-MA) is a selective PI3K inhibitor for Vps34 and PI3Ky with IC50 of 25 µM and 60 µM in HeLa cells; blocks class I PI3K consistently, whereas suppression of class III PI3K is transient, and also blocks autophagosome formation.

Page 8

S2775 Nocodazole

Autophagy Nocodazole is a rapidly-reversible inhibitor of microtubule polymeriza -tion, and also inhibits Abl, Abl(E255K) and Abl(T315I) with IC50 of 0.21 µM, 0.53 µM and 0.64 µM in cell-free assays, respectively. ----- Page 76

S4157 Chloroquine Phosphate

Chloroquine phosphate is a 4-aminoquinoline anti-malarial and anti-rheumatoid agent, also acting as an ATM activator.

----- Page 16

S4430 Hydroxychloroquine Sulfate

Hydroxychloroquine Sulfate is an antimalarial agent used for the treatment of systemic lupus erythematosus, rheumatoid arthritis and other autoimmune, inflammatory and dermatologic conditions. Also acts as an inhibitor of autophagy and toll-like receptor (TLR) 7/9.

Size 10 mg 50 mg 200 mg

S7885 SBI-0206965

SBI-0206965 is a highly selective autophagy kinase ULK1 inhibitor with IC50 of 108 nM, about 7-fold selectivity over ULK2. Size 5 mg 25 mg



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60

Size

Spautin-1 is a potent and specific autophagy inhibitor, and inhibits the deubiquitinating activity of USP10 and USP13 with IC₅₀ of ~0.6-0.7 µM. Size 10 mg 50 mg

Autophagy Activators

S1237 Temozolomide

Temozolomide is a monofunctional SN-1 alkylating agent that can modify nitrogen atoms in the DNA ring and the extracyclic oxygen group, chemically converted to MTIC and degrades to methyldiazonium cation, which transfers methyl groups to DNA at physiologic pH. A DNA damage inducer in L-1210 and L-1210/BCNU cells.

Size 25 mg 100 mg 10 mM/1 mL



Product Citations (4): Nat Med, 2015, 10.1038/nm.3855 Clin Cancer Res, 2014, 20(6): 1555-65

Data from [Clin Cancer Res, 2014, 20(6): 1555-651 Temozolomide (TMZ) purchased from

S1950 Metformin HCI

Metformin HCI decreases hyperglycemia in hepatocytes primarily by suppressing glucose production by the liver (hepatic gluconeogenesis). Size 50 mg 5 g

Product Citations (4): Cancer Cell, 2014, 26(6): 840-50 Oncotarget, 2015, 6(2): 969-78 Data from [Luminescence, 2014, 29(1): 65-73] Metformin HCI (MF) purchased from Selleck

S1047 Vorinostat (SAHA, MK0683)

Vorinostat (suberoylanilide hydroxamic acid, SAHA) is an HDAC inhibitor with IC50 of ~10 nM in a cell-free assay. ----- Page 21

Autophagy S1002 ABT-737

ABT-737 is a BH3 mimetic inhibitor of Bcl-xL, Bcl-2 and Bcl-w with EC50 of 78.7 nM, 30.3 nM and 197.8 nM in cell-free assays, respectively; no inhibition observed against Mcl-1, Bcl-B or Bfl-1. Phase 2. -----Page 57

S1049 Y-27632 2HCI

Y-27632 2HCl is a selective ROCK1 (p160ROCK) inhibitor with Ki of 140 nM in a cell-free assay, exhibiting >200-fold selectivity over other kinases, including PKC, cAMP-dependent protein kinase, MLCK and PAK Page 82

S1039 Rapamycin (Sirolimus) Licensed by Pfizer

Rapamycin (Sirolimus) is a specific mTOR inhibitor with IC50 of ~0.1 nM HEK293 cells. ----- Page 11

S1023 Erlotinib HCI (OSI-744)

Erlotinib HCI (OSI-744) is an EGFR inhibitor with IC50 of 2 nM in cell-free assays, >1000-fold more sensitive for EGFR than for human c-Src or v-Ahĺ Page 39

S1208 Doxorubicin (Adriamycin) Licensed by Pfizer

Doxorubicin (Adriamycin) is an antibiotic agent that inhibits DNA topoisomerase II and induces DNA damage and apoptosis in tumor cells

S1057 Obatoclax Mesylate (GX15-070)

Obatoclax Mesylate (GX15-070) is an antagonist of Bcl-2 with K of 0.22 μM in a cell-free assay, can assist in overcoming MCL-1 mediated resistance to apoptosis. Phase 3.

----- Page 57

S1038 PI-103

PI-103 is a multi-targeted PI3K inhibitor for p110 $\alpha/\beta/\delta/\gamma$ with IC₅₀ of 2 nM/3 nM/3 nM/15 nM in cell-free assays, less potent to mTOR/DNA-PK with IC50 of 30 nM/23 nM. ----- Page 7

S1149 Gemcitabine HCI

Gemcitabine HCl is a DNA synthesis inhibitor with IC50 of 50 nM, 40 nM, 18 nM and 12 nM in PANC1, MIAPaCa2, BxPC3 and Capan2 cells, respectively.

----- Page 88

S2218 Torkinib (PP242)

Torkinib (PP242) is a selective mTOR inhibitor with IC50 of 8 nM in cell-free assays; targets both mTOR complexes with >10- and 100-fold selectivity for mTOR than PI3K δ or PI3K $\alpha/\beta/\gamma$, respectively.

..... Page 11

S1573 Fasudil (HA-1077) HCI

Fasudil (HA-1077), a potent and selective inhibitor of Rho kinase, displays less potent inhibiton over PKA, PKG, PKC and MLCK with K_i of HSN NH NH 1.6, 1.6, 3.3, and 36 µM in cell-free assays, respectively. Page 82

S1972 Tamoxifen Citrate

Tamoxifen Citrate is an antagonist of the estrogen receptor by competitive inhibition of estrogen binding. Page 110

Autophagy Modulators

S1241 Vincristine sulfate

Vincristine sulfate is an inhibitor of polymerization of microtubules by binding to tubulin with IC50 of 32 µM in a cell-free assay. ----- Page 76

S1168 Valproic acid sodium salt (Sodium valproate)

Valproic acid sodium salt (Sodium valproate) is a HDAC inhibitor by selectively inducing proteasomal degradation of HDAC2, used in the treatment of epilepsy, bipolar disorder and prevention of migraine headaches.

----- Page 22

LRRK2 Inhibitor

S7584 LRRK2-IN-1

Excellent Validation, Technical Support and Prompt Delivery

LRRK2-IN-1 is a potent and selective LRRK2 inhibitor with IC50 of 6 nM and 13 nM for LRRK2 (G2019S) and LRRK2 (WT), respectively. Size 10 mg 50 mg 100 mg



JAK/STAT Pathway



JAK Inhibitors

Detailed product information is on page 24-26

Pim Inhibitors

Detailed product information is on page 26

EGFR Inhibitors

Detailed product information is on page 38-40

STAT Inhibitors

innibitory Selectivity							
Inhibitor Name	STAT1	STAT3	STAT5				
S3I-201		+ IC ₅₀ : 86 μM					
Stattic		++ IC50: 5.1 μM					
Niclosamide		+++ IC50: 0.7 μM					
BP-1-102		+++ K _d : 504 nM					
SH-4-54		++++ Kd: 300 nM	++++ Kd: 464 nM				
Cryptotanshinone		++ IC50: 4.6 μM					
Fludarabine	√						
Nifuroxazide	1						
APTSTAT3-9R		1					

of 86 µM in cell-free assays, and low activity towards STAT1 and

0.00

S1155 S3I-201 (NSC 74859)



S3I-201 shows potent inhibition of STAT3 DNA-binding activity with IC50

Notes:

specific value.

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without

JAK / EGFR / Pim / STAT

STAT3 selective



°-∽+⊂

MAPK



MEK Inhibitors

Inhibitory Selectivity

Inhibitor Name	MEK	MEK1	MEK1/2	MEK2	MEK5	Other
Selumetinib		+++ IC50: 14 nM				
PD0325901	++++ IC ₅₀ : 0.33 nM					
Trametinib		++++ IC50: 0.92 nM		++++ IC ₅₀ : 1.8 nM		
U0126-EtOH		+ IC ₅₀ : 0.07 μM		++ IC ₅₀ : 0.06 μM		MKK6/p38 MAPK,MKK3/p38 MAPK
PD184352		++ IC50: 17 nM		++ IC50: 17 nM		
PD98059		+ IC ₅₀ : 2 μM				
BIX 02189					++++ IC50: 1.5 nM	ERK5,TGFβR1
Pimasertib			+ IC ₅₀ : 5 nM-2 μM			
BIX 02188					+++ IC50: 4.3 nM	ERK5,TGFβR1
TAK-733		++++ IC ₅₀ : 3.2 nM				
AZD8330			+++ IC50: 7 nM			ERK phosphorylation
Binimetinib	+++ IC50: 12 nM					
SL-327		+ IC ₅₀ : 0.18 μM		+ IC ₅₀ : 0.22 μM		AP-1,MKK3/p38 MAPK
Refametinib		++ IC ₅₀ : 19 nM		++ IC50: 47 nM		
GDC-0623		++++ IC50: 0.13 nM				
BI-847325		++ IC50: 25 nM		+++ IC50: 4 nM		Aurora B, Aurora C, Aurora A
Cobimetinib		+++ IC ₅₀ : 4.2 nM				
PD318088			N			
Honokiol	1					Akt-phosphorylation

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsss) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

www.selleckchem.com

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

Size 50 mg 1 g 5 g

64

65

MAPK



Raf Inhibitors | Chemical

Inhibitory Selectivity

Inhibitor Name	Raf	C-Raf/Raf-1	B-Raf	Other
Vemurafenib		+ IC50: 48 nM	++ IC50: 100 nM	SRMS,ACK1,MAP4K5 (KHS1)
Sorafenib Tosylate		++++ IC50: 6 nM	++ IC50: 22 nM	VEGFR2/Flk1,mPDGFRβ,PDGFRβ
PLX-4720		+++ IC50: 6.7 nM	+++ IC50: 13 nM	BRK
Dabrafenib		++++ IC50: 5.0 nM	++++ IC50: 0.8 nM	
GDC-0879			++++ IC50: 0.13 nM	
RAF265			++ IC50: 3 nM-60 nM	VEGFR2
AZ 628		++ IC50: 29 nM	++ IC50: 34 nM	
NVP-BHG712		+ IC50: 0.395 μM		EphB4,c-Src,c-Abl
SB590885			++++ K: 0.16 nM	
ZM 336372		+ IC ₅₀ : 70 nM		
Sorafenib		++++ IC50: 6 nM	++ IC50: 38 nM	mVEGFR2(Flk1),mVEGFR3,mPDGFRβ
GW5074		+++ IC50: 9 nM		
TAK-632		++++ IC50: 1.4 nM	+++ IC50: 8.3 nM	Aurora B,PDGFRβ,FGFR3
CEP-32496		++ Kd: 39 nM	+++ K _d : 14 nM	RET,PDGFRβ,LCK
CCT196969		+++ IC50: 0.01 μM	+ IC50: 0.1 μM	LCK,Src,V600E-BRAF
LY3009120		++++ IC50: 4.3 nM	++++ IC50: 5.8~15 nM	
RO5126766		+ IC ₅₀ : 56 nM	+++ IC ₅₀ : 8.2 nM	MEK1
Encorafenib			1	
PLX7904	√			
MLN2480	1			

Notes:

SOX10

respectively.

1. For more details, such as half maximal inhibitory concentrations (ICsss) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation. 3. Red "V" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

Product Citations (62):

Product Citations (38):

4104-151

Selleck

Blood, 2013, 122(9): 1621-33

118-221

Sorafenib Tosylate is a multi-kinase inhibitor of Raf-1, B-Raf and

Nature, 2014, 508(7494); 118-22

Vemurafenib purchased from Selleck

Raf Inhibitors

S1267 Vemurafenib (PLX4032, RG7204)

Size 10 mg 50 mg 200 mg 10 mM/1 mL

Vemurafenib (uM)

A375

Size 5 mg 50 mg 100 mg 10 mM/1 mL

S1040 Sorafenib Tosylate

no VEGF-A VEGF-A

fenib 0 5 10 0 5 10

Vemurafenib (PLX4032, RG7204) is a novel and potent inhibitor of B-Raf^{V600E} with IC₅₀ of 31 nM in cell-free assay. 10-fold selective for B-Raf^{V600E} over wild-type B-Raf in enzymatic assays and the cellular selectivity can exceed 100-fold.



PLX-4720 is a potent and selective inhibitor of B-Raf^{V600E} with IC₅₀ of 13 nM in a cell-free assay, equally potent to c-Raf-1(Y340D and Y341D mutations), 10-fold selectivity for B-RafV600E than for wild-type B-Raf.



S2807 Dabrafenib (GSK2118436)

Dabrafenib (GSK2118436) is a mutant BRAFV600 specific inhibitor with IC50 of 0.8 nM in cell-free assays, with 4- and 6-fold less potency against B-Raf(wt) and c-Raf, respectively



MAPK

Raf

www.selleckchem.com

NO

Dabrafenib purchased from Sellect

Raf / p38 MAPK

S1104 GDC-0879

GDC-0879 is a novel, potent, and selective B-Raf inhibitor with IC50 of 0.13 nM in A375 and Colo205 cells with activity against c-Raf as well; no inhibition known to other protein kinases.

Size 2 mg 10 mg 25 mg 10 mM/1 mL

																1	n—
DMSO	00126	CI-1040	GDC-6879	690680d	DMSO	U0126	CI-1040	GDC-6679	PC69069	DMSO	U0126	CI-1040	GDC-6879	POBACO			Product Citations (13): Cancer Discov, 2013, 3(3): 350-62
-	-	-	-	-	-	-	-	-	-		-	-	÷	-	c-Jun		J Natl Cancer Inst, 2012, 104(21):
		-		-	-	23	1		1	-	1.3	1.9	2.	1.0	P-Erk1/2		1673-9
-	-	=	=	-	=	-	=	=	=	=	=	=	=	=	Erk1/2		
•		-			•	•	-	•	•	-			-		GAPDH		Data from [J Natl Cancer Inst, 2012,
-		VM1	15			LC	ox-	мv		·	н	T-14	44			5	104(21): 1673-9]
																/	CDC 0979 purchased from Selleck

S7108 Encorafenib (LGX818)

Encorafenib (LGX818) is a highly potent RAF inhibitor with selective anti-proliferative and apoptotic activity in cells expressing B-RAF(V600E) with EC50 of 4 nM. Phase 3.

Size 1 mg 5 mg

MAPK S7397 Sorafenib (BAY 43-9006)

Sorafenib is a multi-kinase inhibitor of Raf-1, B-Raf and VEGFR-2 with IC50 of 6 nM, 22 nM and 90 nM in cell-free assays, respectively.

Size 20 mg 50 mg 200 mg





Sorafenib purchased from Selleck

S7291 TAK-632

TAK-632 is a potent pan-Raf inhibitor with IC50 of 8.3 nM and 1.4 nM for B-Raf(wt) and C-Raf in cell-free assays, respectively, showing less or no inhibition against other tested kinases.

Size 5 mg 20 mg

S2746 AZ 628

AZ 628 is a new pan-Raf inhibitor for BRAF, BRAFV600E, and c-Raf-1 with IC50 of 105 nM, 34 nM and 29 nM in cell-free assays, and also inhibits VEGFR2, DDR2, Lyn, Flt1, FMS, etc.

Size 5 mg 25 mg 10 mM/1 mL



SB590885 is a potent B-Raf inhibitor with K of 0.16 nM in a cell-free assay, 11-fold greater selectivity for B-Raf over c-Raf, no inhibition to other human kinases.

Size 10 mg 50 mg 10 mM/1 mL





15-0---



Raf Chemical

S2872 GW5074

B-Raf selective

B-Raf selective

aray

Size

In Na

SE

Do

S7842 LY3009120

LY03009120 is a potent pan-Raf inhibitor with IC50 of 44 nM, 31-47 nM, and 42 nM for A-raf, B-Raf, and C-Raf in A375 cells, respectively. Phase

GW5074 is a potent and selective c-Raf inhibitor with IC50 of 9 nM, but

Jan Hink

C-Raf/Raf-1 selective

p38 MAPK Inhibitors

Inhibitory Selectivity

5 mg 25 mg 100 mg

hibitor ame	p38 MAPK	p38α	p38β	Other
203580	+ IC50: 0.3-0.5 μM			PKB
ramapimod		++++Kd: 0.1 nM		
202190		++ IC50: 50 nM	++ IC50: 100 nM	
2228820		++++ IC50: 7 nM		
-702		+++ IC50: 4-20 nM		
-797804		+++ IC ₅₀ : 26 nM	+ IC ₅₀ : 102 nM	
-745		+++ IC50: 10 nM	+ IC50: 220 nM	
K-715		++++IC50: 7.1 nM	+ IC50: 0.20 μM	
mapimod		+++ IC ₅₀ : 0.014 μM	+ IC ₅₀ : 0.48 μM	
239063		++ IC50: 44 nM	++ IC50: 44 nM	
smapimod		+++ pKi: 8.1	+++ pKi: 7.6	
epinone-L		++++IC50: 5 nM		
xmetinib	1			Tie-2

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

S1076 SB203580 (RWJ 64809)

SB203580 is a p38 MAPK inhibitor with IC $_{50}$ of 0.3-0.5 μM in THP-1 cells, 10-fold less sensitive to SAPK3(106T) and SAPK4(106T) and blocks PKB phosphorylation with IC50 of 3-5 µM.



S7215 Losmapimod (GW856553X, GW856553, GSK-AHAB)

Losmapimod (GW856553X) is a selective, potent, and orally active p38 MAPK inhibitor with pK of 8.1 and 7.6 for p38a and p38b, respectively. Phase 3. Size 10 mg 50 mg



S1574 Doramapimod (BIRB 796)

Doramapimod (BIRB 796) is a pan-p38 MAPK inhibitor with IC50 of 38 nM, 65 nM, 200 nM and 520 nM for p38α/β/γ/δ in cell-free assays, and binds p38α with Kd of 0.1 nM in THP-1 cells, 330-fold greater selectivity versus JNK2, weak inhibition for c-RAF, Fyn and Lck, insignificant inhibition of ERK-1, SYK, IKK2,



S1494 LY2228820

LY2228820 is a novel and potent inhibitor of p38 MAPK with IC50 of 7 nM, but does not alter p38 MAPK activation. Phase 1/2.

Size 5 mg 10 mg 50 mg 10 mM/1 mL



Blood, 2012, 119(26): 6255-8 Data from [Blood, 2012, 119(26) LY2228820 purchased from Selleck

S1077 SB202190 (FHPI) p38a selective

SB202190 (FHPI) is a potent p38 MAPK inhibitor targeting p38α/β with IC50 of 50 nM/100 nM in cell-free assays, sometimes used instead of SB 203580 to investigate potential roles for SAPK2a/p38 in vivo. Size 25 mg 100 mg 10 mM/1 mL



S6005 VX-702

n38a selective

VX-702 is a highly selective inhibitor of p38a MAPK, 14-fold higher potency against the p38α versus p38β. Phase 2.



S8125 Pamapimod (R-1503, Ro4402257)

Pamapimod (R-1503, Ro4402257) is a novel, selective inhibitor of p38 mitogen-activated protein kinase. It inhibits p38a and p38β enzymatic activity with IC50 values of 0.014 ± 0.002 and 0.48 ± 0.04 microM, respectively with no activity against p38delta or p38gamma isoforms.





p38a selective

MAPK

JNK Inhibitors

Inhibitory Selectivity

Inhibitor Name	JNK1	JNK2	JNK3	JNK	Other
SP600125	+++ ICso: 40 nM	+++ IC50: 40 nM	++ IC50: 90 nM	+ ICso: 0.4 μM	Aurora A, TrkA, FLT3
JNK-IN-8	++++ IC50: 4.7 nM	+++ IC50: 18.7 nM	++++ IC50: 1 nM		Kit (V559D,T670I),Kit (V559D),RIOK2
JNK Inhibitor IX		+ pIC50: 6.5	++ pIC50: 6.7		

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsss) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

S1460 SP600125 (Nsc75890)

SP600125 is a broad-spectrum JNK inhibitor for JNK1, JNK2 and JNK3 with IC50 of 40 nM, 40 nM and 90 nM in cell-free assays, respectively; 10-fold greater selectivity against MKK4; 25-fold greater selectivity against MKK3, MKK6, PKB, and PKCa, and 100-fold selectivity against ERK2, p38, Chk1, EGFR etc.

Size 10 mg 50 mg 200 mg 10 mM/1 mL



S4901 JNK-IN-8

JNK-IN-8 is the first irreversible JNK inhibitor for JNK1, JNK2 and JNK4 with IC50 of 4.7 nM, 18.7 nM and 1 nM, >10-fold selectivity against MNK2, Fms and no inhibition to c-Kit, Met, PDGFRβ in A375 cell line. Size 5 mg



JNK-IN-8 purchased from Selleci

32(4): 626-351 SB590885 purchased from Selleck

68

Data from [J Neurosci, 2013, 33(7);

Sk

JNK / ERK / MNK

S7409	Anisomycin
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Anisomycin is an antibiotic, which inhibits protein synthesis, and also acts as a JNK activator.

Size 10 mg 50 mg 200 mg

500

ERK Inhibitors

Inhibitory Selectivity

Inhibitor Name	ERK1	ERK2	ERK5	ERK	Other
SCH772984	+++ IC ₅₀ : 4 nM	++++ IC ₅₀ : 1 nM			
LY3214996	+++ ICso: 5 nM	+++			
SC1	++ Kd:98 nM				RasGAP
VX-11e		+++ Ki: <2 nM			
DEL-22379			+ IC ₅₀ : 0.5 μM	+ IC ₅₀ : 0.5 μM	
Ulixertinib		++++ ICso: <0.3 nM			
GDC-0994	+++ ICso: 1.1 nM	++++ ICso: 0.3 nM			
FR 180204	+ Κ _i : 0.31 μΜ	++ Κi: 0.14 μM			
ERK5-IN-1			++ IC ₅₀ : 162 nM		

Notes:

MAPK

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

S7101 SCH772984

Size 5 mg

in RAS- or BRAF-mutant cancer cells.

S7554 GDC-0994

SCH772984 is a novel, specific inhibitor of ERK1/2 with IC₅₀ values of 4 GDC-0994 is a potent, orally available and highly selective ERK1/2 nM and 1 nM in cell-free assay, respectively, And show robust efficacy inhibitor with IC50 of 1.1 nM and 0.3 nM, respectively. Phase 1. 5 mg ____ 25 mg



S7854 Ulixertinib (BVD-523, VRT752271)

Ulixertinib (BVD-523, VRT752271) is a potent and reversible ERK1/ERK2 inhibitor with IC50 of <0.3 nM for ERK2. Phase 1 5 mg 25 mg 100 mg Size

S7525 XMD8-92

XMD8-92 is a potent and selective BMK1/ERK5 inhibitor with K_d of 80 nM.

Product Citations (7): J Clin Invest, 2015, 125(6): 2484-96

1-8]

Cell Res, 2015, 10.1038/cr.2015.30

SCH772984 purchased from Selleck

Data from [Leuk Lymphoma, 2014, 22:

ERK5 selective



S7524 FR 180204

FR 180204 is an ATP-competitive, selective ERK inhibitor with K_i of 0.31 µM and 0.14 µM for ERK1 And ERK2, respectively. It is 30-fold less potent against the related kinase p38a and failed to inhibit any kinases(MEK1, MKK4, IKKa, PKCa, Src, Syc, and PDGFa) at less than 30 µM.

Size 5 mg 25 mg

NN-N N NH, 050

MNK Inhibitor	
S8275 eFT-508 (eFT508)	
eFT-508 (eFT508) is a potent and selective MNK1/2 inhibitit of 2.4 nM and 1 nM, respectively. It potentially results in tumor cell proliferation and tumor growth.	or with IC50s decreased
Size 2 mg 5 mg 25 mg	wate

Cytoskeletal Signaling



Akt Inhibitors

Detailed product information is on page 12-13

Bcr-Abl Inhibitors

Detailed product information is on page 51-52

FAK Inhibitors

Detailed product information is on page 54

Wnt/beta-catenin Inhibitors

Inh	ibito	rv Se	lectivity

Inhibitor Name	Wnt/beta-catenin	Other
XAV-939	+++ IC ₅₀ : 11 nM	
ICG-001	+ IC ₅₀ : 3 μM	
IWR-1-endo	+ IC50: 180 nM	
Wnt-C59	++++ IC50: 74 pM	
IWP-2	++ IC ₅₀ : 27 nM	
IWP-L6	++++ EC50: 0.5 nM	
KYA1797K	+ ICso: 0.75 μM	
PRI-724	++ IC50: 150 nM	
WIKI4	+++ IC ₅₀ : 15 nM	
LGK-974	1	
KY02111	1	
FH535	1	PPARγ,PPARδ

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

S1180 XAV-939

XAV-939 selectively inhibits Wnt/β-catenin-mediated transcription through tankyrase1/2 inhibition with IC50 of 11 nM/4 nM in cell-free assays, regulates axin levels and does not affect CRE, NF-kB or TGF-β.

Size 50 mg 200 mg 10 mM/1 mL Cytoskeletal Signaling 10 mg A3-0-* . Product Citations (21): Nat Cell Biol, 2014, 16(2): 179-90 Nat Commun. 2014. 5: 5455 Data from [J Mol Cell Cardiol, 2013, 62: 203-13] XAV-939 purchased from Selleck

S2662 ICG-001

ICG-001:

Survivin

Tubulin

ICG-001 antagonizes Wnt/β -catenin/TCF-mediated transcription and specifically binds to CREB-binding protein (CBP) with IC50 of 3 µM, but is not the related transcriptional coactivator p300.



 $\mathrm{Class}_{\mathrm{H}}$ \mathbb{Q} Product Citations (13): AsPC-1 L3.6pl PANC-1 MiaPaCa-2 Proc Natl Acad Sci USA. 2013, 110(52): E5039-48



S7143 LGK-974

LGK-974 is a potent and specific PORCN inhibitor, and inhibits Wnt signaling with IC50 of 0.4 nM in TM3 cells. Phase 1.



PKC Inhibitors

Inhibitory Selectivity

Inhibitor Name	РКС	ΡΚCα	рксβ	РКСү	PKCδ	ΡΚCε	ΡΚϹζ	РКСη	РКСӨ	РКСµ	Other
Enzastaurin		++ IC ₅₀ : 39 nM	+++ IC50: 6 nM	+ IC ₅₀ : 83 nM		+ IC ₅₀ : 110 nM					
Sotrastaurin		++++ K _i : 0.95 nM	+++++ K _i : 0.64 nM		+++++ K _i : 2.1 nM	+++++ K _i : 3.2 nM		+++++ K _i : 1.8 nM	++++ Ki: 0.22 nM		
Staurosporine		++++ IC ₅₀ : 2 nM		++++ IC ₅₀ : 5 nM	++ IC ₅₀ : 20 nM	+ IC ₅₀ : 73 nM	+ IC ₅₀ : 1086 nM	++++ IC ₅₀ : 4 nM			c-Fgr,phosphorylase kinase,S6 kinase
Go 6983		+++ IC ₅₀ : 7 nM	+++ IC ₅₀ : 7 nM	+++ IC ₅₀ : 6 nM	+++ IC ₅₀ : 10 nM		++ IC ₅₀ : 60 nM			+ IC ₅₀ : 20 μM	
Bisindolylmaleimide I		++ IC ₅₀ : 20 nM	+++ IC ₅₀ : 17 nM	++ IC ₅₀ : 20 nM							PDGFR
Ro 31-8220 Mesylate		++++ IC ₅₀ : 5 nM	+++ IC ₅₀ : 24 nM	++ IC ₅₀ : 27 nM		++ IC ₅₀ : 24 nM					
Dequalinium Chloride	+ IC50: 7-18 μM										
Midostaurin		++ IC50: 22 nM	++ IC50: 30 nM	++ IC50: 24 nM	+ IC50: 330 nM	+ IC50: 1.25 μM	+ IC50: 465 μM	+ IC50: 160 nM			PPK,KDR,c-Syk
Go6976	+++ IC50: 7.9 nM	++++ IC50: 2.3 nM	+++ IC50: 6.2 nM								FLT3,JAK2

S7086 IWR-1-endo

IWR-1-endo is a Wnt pathway inhibitor with IC50 of 180 nM in L-cells expressing Wnt3A, induces Axin2 protein levels and promotes β-catenin phosphorylation by stabilizing Axin-scaffolded destruction complexes. Size 10 mg 25 mg

S7037 Wnt-C59 (C59)

Wnt-C59 (C59) is a PORCN inhibitor for Wnt3A-mediated activation of a multimerized TCF-binding site driving luciferase with IC50 of 74 pM in HEK293 cells. Size 5 mg

Dauo⁰

S7085 IWP-2

IWP-2 is an inhibitor of Wnt processing and secretion with IC50 of 27 nM in a cell-free assay, selective blockage of Porcn-mediated Wnt palmitoylation, does not affect Wnt/β-catenin in general and displays no effect against Wnt-stimulated cellular responses. Size 10 mg 50 mg

S7096 KY02111

S8262 PRI-724

KY02111 promotes differentiation of hPSCs to cardiomyocytes by inhibiting Wnt signaling, may act downstream of APC and GSK3β. Size 10 mg 50 mg

PRI-724 is a potent, specific inhibitor of the canonical Wnt signaling pathway in cancer stem cells with potential antineoplastic activity. PRI-724 specifically inhibits the recruiting of beta-catenin with its coactivator CBP. Size 5 mg 25 mg 100 mg

Inhibitory Selectivity

Inhibitor Name	РКС	ΡΚCα	рксβ	РКСү	PKCδ	ΡΚϹε	РКСζ	ΡΚϹη	РКСӨ	ΡΚϹμ	Other
Quercetin	V										Sirtuin,Src,PI3Ky
Myricitrin		۸									

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsss) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

S1055 Enzastaurin (LY317615)

Enzastaurin (LY317615) is a potent PKCβ selective inhibitor with IC50 of 6 nM in cell-free assays, 6- to 20-fold selectivity against PKCa, PKCy and PKC₂. Phase 3.

Size 10 mg 50 mg 200 mg 10 mM/1 mL



S2791 Sotrastaurin (AEB071)

Sotrastaurin is a potent and selective pan-PKC inhibitor, mostly for PKC θ with K of 0.22 nM in a cell-free assay; inactive to PKC ζ . Phase 2.



Product Citations (4): Proc Natl Acad Sci USA, 2014, 111(15): E1528-37 Cancer Cell, 2015, 27(3): 397-408 Data from [Proc Natl Acad Sci USA. 2014, 111(15): E1528-37] Sotrastaurin (So) purchased from

S1421 Staurosporine (CGP 41251)

Staurosporine is a potent PKC inhibitor for PKCq, PKCq and PKCq with IC50 of 2 nM, 5 nM and 4 nM, less potent to PKCo (20 nM), PKCE (73 nM) and little active to PKCZ (1086 nM) in cell-free assays. Staurosporine also shows inhibitory activities on other kinases, such as PKA, PKG, S6K, CaMKII etc. Phase 3. N.o

Size 2 mg



Product Citations (5): Cancer Res, 2014, 74(23): 7090-102 J Biomol Screen, 2013, 18(4): 388-99 Data from [J Biomol Screen, 2013,

020

18(4): 388-99] Staurosporine purchased from Selleck



Cytoskeletal Signaling Go 6983 is a pan-PKC inhibitor against for PKCa, PKCβ, PKCγ and PKCō with IC50 of 7 nM, 7 nM, 6 nM and 10 nM, respectively; less potent to PKCζ and inactive to PKCμ.



S7208 BisindolyImaleimide I (GF109203X)

GF109203X is a potent PKC inhibitor with IC50 of 20 nM, 17 nM, 16 nM, and 20 nM for PKCa, PKCBI, PKCBII, and PKCy, respectively, showing more than 3000-fold selectivity for PKC as compared to EGFR, PDGFR and insulin receptor.



S7119 Go6976

Size 5 mg 25 mg

1 mg 10 mg

Size

Go6976 is a potent PKC inhibitor with IC50 of 7.9 nM, 2.3 nM, and 6.2 nM for PKC (Rat brain), PKCa, and PKCB1, respectively. Also a potent inhibitor of JAK2 and Elt3



S7207 Ro 31-8220 Mesylate (Bisindolylmaleimide IX Mesylate)

Ro 31-8220 Mesylate is a pan-PKC inhibitor with IC50 of 5 nM, 24 nM, 14 nM, 27 nM, and 24 nM for PKC-a, PKC-BI, PKC-BI, PKC-y, and PKC-ɛ, respectively, and also shows potent inhibition against MAPKAP-K1b, MSK1, GSK3ß and S6K1.

Size 10 mg 50 mg

Quercetin, a natural flavonoid present in vegetables, fruit and wine, is a stimulator of recombinant SIRT1 and also a PI3K inhibitor with IC50 of 2.4-5.4 uM. Phase 4. ----- Page 30

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Product Citation (1): BMC Genomics, 2014, 15(1): 263

15(1): 263 1

Data from [BMC Genomics, 2014,

Elesciomol purchased from Selleck

HSP (e.g. HSP90) Modulator

elicits pro-apoptosis events among tumor cells. Phase 3.

elesciomol 0.228 µN

5 mg 10 mg 50 mg 10 mM/1 mL

control disulfiram 0.228 µM

Elesclomol (STA-4783) is a novel potent oxidative stress inducer that

HSP (e.g. HSP90) Inhibitors | Modulator

Inhibitory Selectivity

Inhibitor Name	HSP70	HSP90	HSP90α	HSP90ß	HSP105	Other
Tanespimycin		+++ IC50: 5 nM				
Luminespib		+++ IC50: 13 nM	+++ IC50: 13 nM	+++ IC50: 21 nM		
Alvespimycin HCI		+ IC50: 62 nM				
Ganetespib		+++ IC50: 4 nM				
BIIB021		++++ EC50: 38 nM				
Onalespib		+++ IC50: 18 nM				
Geldanamycin		+ Kd: 1.2 μΜ				p185
NVP-BEP800		+ IC50: 58 nM		+ IC ₅₀ : 58 nM		
SNX-2112		++ K _a : 30 nM	++ Ka: 30 nM	++ Ka: 30 nM		
PF-04929113		++ Kd: 41 nM				HER2
KW-2478		++++ IC50: 3.8 nM				
XL888		++ IC50: 24 nM				
Apoptozole	+ ICso: 0.14 μM					
VER155008	+ IC ₅₀ : 0.5 μM					
VER-50589		+++ IC50: 21 nM		+++ ICso: 21 nM		
CH5138303		++++ K _d : 0.48 nM	++++ K _d : 0.48 nM			
VER-49009		++ IC50: 47 nM		++ IC ₅₀ : 47 nM		
NMS-E973		+++ DC50: <10 nM				
PU-H71		+ ICso: 51 nM				
HSP990		++++ IC50: 0.8 nM	++++ IC50: 0.6 nM	++++ IC50: 0.8 nM		
KNK437					√	

Cytoskeletal Signaling

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

HSP (e.g. HSP90) Inhibitors

S1141 Tanespimycin (17-AAG)

Tanespimycin (17-AAG) is a potent HSP90 inhibitor with IC50 of 5 nM in a cell-free assay, having a 100-fold higher binding affinity for HSP90 derived from tumor cells than for HSP90 from normal cells. Phase 2.

Size 25 mg 100 mg 10 mM/1 mL

74



BIIB021 is an orally available, fully synthetic small-molecule inhibitor of HSP90 with K₁ and EC₅₀ of 1.7 nM and 38 nM, respectively. Phase 2.



AUY-922 purchased from Selleck S1159 Ganetespib (STA-9090) Ganetespib (STA-9090) is an HSP90 inhibitor with IC50 of 4 nM in OSA 8 cells, inducing apoptosis of OSA cells while normal osteoblasts are not affected; active metabolite of STA-1474. Phase 3. Size 5 mg 10 mg 10 mM/1 mL

S1069 Luminespib (AUY-922, NVP-AUY922)

Luminespib (AUY-922, NVP-AUY922) is a highly potent HSP90 inhibitor

for HSP90α/β with IC50 of 13 nM /21 nM in cell-free assays, weaker

potency against the HSP90 family members GRP94 and TRAP-1,

exhibiting the tightest binding of any small-molecule HSP90 ligand.

S1142 Alvespimycin (17-DMAG) HCI

Alvespimycin (17-DMAG) HCl is a potent HSP90 inhibitor with IC50 of 62 nM in a cell-free assay. Phase 2.

Size 25 mg 100 mg 200 mg 10 mM/1 mL



S1163 Onalespib (AT13387)

Onalespib (AT13387) is a selective potent Hsp90 inhibitor with IC50 of 18 nM in A375 cells, displaying a long duration of anti-tumor activity. Phase 2.



Data from [PLoS One, 2013, 8(4): 59315] AT13387 purchased from Selleck

Product Citation (1): PLoS One, 2013, 8(4): e59315

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S8039 PU-H71 (NSC 750424)

PU-H71 is a potent and selective inhibitor of HSP90 with IC50 of 51 nM. Phase 1. ĠЪ В Size 10 mg 25 mg

S2713 Geldanamycin

Geldanamycin is a natural existing HSP90 inhibitor with Kd of 1.2 µM. specifically disrupting glucocorticoid receptor (GR)/HSP association.

Size 5 mg 10 mg 50 mg 10 mM/1 mL

S7751 VER155008

VER155008 is a potent Hsp70 family inhibitor with IC50 of 0.5 µM, 2.6 µM, and 2.6 µM in cell-free assays for HSP70, HSC70, and GRP78, respectively, >100-fold selectivity over HSP90.

Size 10 mg 50 mg

S7097 HSP990 (NVP-HSP990)

NVP-HSP990 (HSP990) is a novel, potent and selective HSP90 inhibitor for HSP90a/B with IC50 of 0.6 nM/0.8 nM.

Size 5 mg 25 mg 100 mg





Product Citations (2): J Biol Chem, 2013, 288(49): 35149-58 Oncogenesis, 2014, 3: e100

AZ 3146 purchased from Selleck

 $\mathbb{Q}_{q^kg} \mathfrak{U}_g \mathfrak{U}_g \mathbb{Q}$

Kinesin Inhibitors

Inhibitory Selectivity

S1052 Elesciomol (STA-4783)

11.6

BCS [µM]

Size

	-
Inhibitor Name	Kinesin
Ispinesib	+++ K _i app: 1.7 nM
SB743921	++++ ICso: 14.4 nM
AZ 3146	+ ICso: ~35 nM
GSK923295	++ Ki: 3.2 nM
MPI-0479605	+++ IC ₅₀ : 1.8 nM
ARQ 621	1
Notes:	

1. For more details, such as half maximal inhibitory concentrations (IC50s) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

S1452 Ispinesib (SB-715992, CK0238273)

Ispinesib (SB-715992) is a potent, specific and reversible inhibitor of kinesin spindle protein (KSP) with Ki app of 1.7 nM in a cell-free assay, no inhibition to CENP-E, RabK6, MCAK, MKLP1, KHC or Kif1A. Phase 2.

recruitment of CENP-E (kinesin-related motor protein), less potent to

10 mg 50 mg 10 mM/1 mL Size



-> MCF7 (Lu)

- T47D (Lu) - ZR751 (Lu)

HCC1143 (Ba

MDA468 (Ba)
 MDA435 (Ba)
 MDA436 (Ba)
 BT20 (Ba)
 MDA231 (Ba)

Nat Methods, 2015, 10.1038/nmeth.3363 Nat Commun. 2015

Data from [Mol Oncol, 2014, pii: S1574-7891(14)00131-8] Ispinesib purchased from Sellect

Data from [Oncogenesis, 2014, 3: e1001

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S7090 GSK923295

GSK923295 is a first-in-class, specific allosteric inhibitor of CENP-E kinesin motor ATPase with K of 3.2 nM, and less potent to mutant 1182 and T183. Phase 1.

Size 5 mg 50 mg

Microtubule Associated Inhibitors, **Inhibitory Selectivity**

Inhibitor Name	Microtubule Associated	Other	microtubules by binding to stabilize
	microtubule Associated	other	Size 10 mg 50 mg 10 mM/1 mL
Paclitaxel	++++ IC50: 0.1 pM		
Vincristine sulfate	+ IC50: 32 μM		
Patupilone	+++ EC0.01: 1.8 μM		- <u>Docetaxel</u> <u>Thiostrepton</u> <u>Co-treat</u> 0h 24h 48h 72h (h 24h 48h 72h 0h 24h 48h 72h 0h 24h 48h 72h
Lexibulin (CYT997)	++++ IC50: 10-100 nM		P0001
Epothilone A	++ EC0.01: 2 μM		UST MI
Fosbretabulin Disodium	++ IC50: 2.4 μM		
Vinflunine Tartrate	+++ IC50: 1.2 μM		S1364 Patupilone (EPO906, Epothi
CW069	+ IC50: 75 μM		Patupilone (EPO906, Epothilone
Combretastatin A4	+++ Kd: 0.4 μM		stabilizing agent with EC0.01 of 1.8 µ
CK-636	++ IC50: 4 μM		Size 2 mg 10 mg 25 mg 10
Docetaxel	1		100
ABT-751 (E7010)	1		80-
Nocodazole	V	Abl,Abl (E255K),Abl (T315I)	
Cabazitaxel	1		S 40- + MCF10A
Vinblastine	V	nAChR	** 20-
Albendazole	1		
Docetaxel Trihydrate	1		1 10 100 1000 10000 100000 Epothilone B (nM)
TAI-1	1		
INH6	1		S4269 Vinorelbine Lartrate
INH1	1		mitosis through interaction with tub
Vinorelbine Tartrate	√		Size 10 mg 25 mg 50 mg
Triclabendazole	1		
Orless Arbeits	1		Energi of Vinoreibine on NSCLC cell lines

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without

specific value

S1241 Vincristine sulfate

Vincristine sulfate is an inhibitor of polymerization of microtubules by binding to tubulin with IC50 of 32 µM in a cell-free assay.







S1148 Docetaxel (RP56976, NSC 628503)

Docetaxel, an analog of taxol, is an inhibitor of depolymerisation of nicrotubules by binding to stabilized microtubules. ize 10 mg 50 mg 10 mM/1 mL Product Citations (7): Int J Cancer, 2015, 136(9): 2065-77 - Docetaxel Thiostrepton Co-treated Biochem Pharmacol, 2015 10.1016/j.bcp.2015.05.005

> Data from [J Transl Med. 2013, 11: 204] Docetaxel purchased from Selleck

S1364 Patupilone (EPO906, Epothilone B)

Patupilone (EPO906, Epothilone B) is a paclitaxel-like microtubuletabilizing agent with EC0.01 of 1.8 µM. Phase 2.



4269 Vinorelbine Tartrate

/inorelbine Tartrate is a semi-synthetic vinca alkaloid, and inhibits nitosis through interaction with tubulin.



S2775 Nocodazole

Nocodazole is a rapidly-reversible inhibitor of microtubule polymerization, and also inhibits Abl, Abl(E255K) and Abl(T315I) with IC50 of 0.21 µM, 0.53 µM and 0.64 µM in cell-free assays, respectively. Size 10 mg 50 mg

S4505 Vinblastine sulfate

5 mg 25 mg 100 mg

Vinblastine sulfate inhibits microtubule formation and suppresses nAChR activity with IC50 of 8.9 µM in a cell-free assay, used to treat certain kinds of cancer



Microtubule Associated / Integrin / PAK / Dynamin

S7094 PF-3758309 (PF-03758309) Fosbretabulin (Combretastatin A4 Phosphate (CA4P)) Disodium is the

PF-03758309 is a potent, ATP-competitive, pyrrolopyrazole inhibitor of PAK4 with IC50 of 1.3 nM

Size 10 mg 50 mg



FRAX597 is a potent, ATP-competitive inhibitor of group IPAKs with IC50 of 8 nM, 13 nM, and 19 nM for PAK1, PAK2, and PAK3, respectively.

5 mg 25 mg gande

Dynamin Inhibitors

Inhibitory Selectivity

Inhibitor Name	Dynamin
Dynasore	++ ICso: ~15 μM
Mdivi-1	+++ IC ₅₀ : 1-10 μM
Dyngo-4a	++++ IC ₅₀ : 0.38 μM

Cilengitide is a potent integrin inhibitor for $\alpha\nu\beta3$ receptor and $\alpha\nu\beta5$ Notes: receptor with IC50 of 4.1 nM and 79 nM, respectively; ~10-fold selectivity

1. For more details, such as half maximal inhibitory concentrations (IC50s) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

S8047 Dynasore

Dynasore is a cell-permeable, reversible non-competitive dynamin inhibitor of GTPase activity of dynamin 1/2, with IC50 of 15 µM in a cell-free assay, and also inhibits the mitochondrial dynamin Drp1, with no effect against other small GTPase.

Size 10 mg 50 mg

Product Citation (1): PLoS One, 2014, 9(4); e94732 Data from [PLoS One, 2014, 9(4): e94732] Dynasore purchased from Selleck

integrin $\alpha 5\beta 1$. It binds to several integrins, including $\alpha 5\beta 1$ and $\alpha \nu \beta 3$,

PAK Inhibitors

S7093 IPA-3

Mdivi-1 is a selective cell-permeable inhibitor of mitochondrial division DRP1 (dynamin-related GTPase) and mitochondrial division Dynamin I (Dnm1) with IC50 of 1-10 µM. Size 20 mg 50 mg

S7163 Dyngo-4a

Dvngo-4a is a potent dvnamin inhibitor with IC50 of 0.38 µM, 1.1µM, and 2.3 µM for Dynl (brain), Dynl (rec), and Dynll (rec), respectively. Size 10 mg 50 mg 200 mg



IPA-3 is a selective non-ATP competitive Pak1 inhibitor with IC50 of 2.5 µM, no inhibition to group II PAKs (PAKs 4-6). Size 5 mg 50 mg

www.selleckchem.com





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Size

WALCH ,



against gpllbllla. Phase 2.

Integrin Inhibitors

S7077 Cilengitide (EMD 121974, NSC 707544)

S8008 RGD (Arg-Gly-Asp) Peptides

RGD (Arg-Gly-Asp) Peptides is a cell adhesion motif which can mimic cell adhesion proteins and bind to integrins.

S7204 Fosbretabulin (Combretastatin A4 Phosphate (CA4P)) Disodium

water-soluble prodrug of Combretastatin A4 (CA4), which is a

microtubule-targeting agent that binds β -tubulin with Kd of 0.4 μ M in a

cell-free assay. Fosbretabulin Disodium inhibits the polymerization of

tubulin with IC50 of 2.4 µM, and also disrupts tumor vasculature. Phase 3.

Ixabepilone is an orally bioavailable microtubule inhibitor. It binds to

tubulin and promotes tubulin polymerization and microtubule

stabilization, thereby arresting cells in the G2-M phase of the cell cycle

Integrin Inhibitors | Antagonist

Size 10 mg 25 mg

Size 5 mg

S7930 Ixabepilone (BMS-247550)

and inducing tumor cell apoptosis.

Size 10 mg

Integrin Antagonist

S8454 ATN-161 (Ac-PHSCN-NH2)

ATN-161 (Ac-PHSCN-NH2) is a novel small peptide antagonist of that play a role in angiogenesis and tumor progression.

Size 5 mg 25 mg 100 mg



Aurora Kinase / CDK

Cell Cycle



Aurora Kinase Inhibitors

Detailed product information is on page 27-29

CDK Inhibitors

Inhibitory Selectivity

Inhibitor Name	CDK1	CDK2	CDK3	CDK4	CDK5	CDK6	CDK7	CDK9	CLK	CDK	Cdc	Other
Palbociclib HCI				+++++ IC50: 11 nM		+++ IC50: 15 nM						
Roscovitine		+ ICso: 0.7 μM			++ ICso: 0.16 μM						+ ICso: 0.65 μM	ERK2,GST-ERK1,ERK1
SNS-032	+ IC50: 480 nM	+++ IC50: 38 nM		+ IC50: 925 nM	+ IC50: 340 nM		++ IC50: 62 nM	+++++ IC50: 4 nM				GSK-3α,GSK-3β
Dinaciclib	+++++ IC ₅₀ : 3 nM	+++++ IC ₅₀ : 1 nM			+++++ IC ₅₀ : 1 nM			+++++ IC ₅₀ : 4 nM				
Flavopiridol	+++ IC ₅₀ : 40 nM	+++ IC ₅₀ : 40 nM		+++ IC ₅₀ : 40 nM		+++ IC ₅₀ : 40 nM	+ IC ₅₀ : 300 nM					

				IC 50: U. 16 µM	
Pan-Cnk AZD Selective Ch	Innibitor 7762 k Inhibitors 8 (Chk1)	PHA-793887	++ IC ₅₀ : 60 nM	++++ IC ₅₀ : 8 nM	
MK-8776 (Chk1) CHIR-124 (Chk1)		BS-181 HCI			
		Palbociclib Isethionate			
Cyclin-H		A-674563		++ Ki: 46 nM	
		abemaciclib			
-		BMS-265246	++++ IC ₅₀ : 6 nM	++++ IC ₅₀ : 9 nM	
		PHA-767491	++ IC50: 250 nM	++ IC50: 240 nM	
		Milciclib	+ IC50: 398 nM	+++ IC50: 363 nM	
n-A		R547	++++ Ki: 2 nM	++++ K:: 3 nM	
		NU6027	+ Κι: 2.5 μΜ	+ Κ;: 1.3 μΜ	
ar Exclusion		P276-00	++ IC ₅₀ : 79 nM	++ IC50: 224 nM	
		Kenpaullone	+ ICso: 0.4μM	+ IC50: 0.68µM	
53 Activators		K03861		++++ Kd: 15.4 nM	
NSC 319726 53 Inhibitors Pifithrin-a		THZ1 2HCI			
Pifithrin-µ		AT7519 HCI	++ IC50: 210 nM	++ IC50: 47 nM	+ IC ₅₀ : 3
Idm2 Antagonists Nutlin-3a Nutlin-3a		Purvalanol A		+++ IC ₅₀ : 70 nM	
YH239-EE Mdm2Activator NSC 207895		Ro-3306	+++ Ki: 20 nM		
		SU9516	+++ IC50: 40 nM	+++ IC50: 22 nM	
		XL413			
NU-7441 NU7026 VIL 006649		LDC000067	+ IC ₅₀ : 5.513 μΝ	+ IC50: 2.441 μM	
PIK-75		ML167			

Inhibitor Name	CDK1	CDK2	CDK3	CDK4	CDK5	CDK6	CDK7	CDK9	CLK	CDK	Cdc	Other
AT7519	++ IC ₅₀ : 210 nM	++ IC ₅₀ : 47 nM	+ IC ₅₀ : 360 nM	++ IC ₅₀ : 100 nM	+++ IC ₅₀ : 13 nM	++ IC ₅₀ : 170 nM	+ IC ₅₀ : 2.4 μM	++++ IC ₅₀ : <10 nM				GSK-3β
Flavopiridol HCl	+++ IC ₅₀ : 40 nM	+++ IC ₅₀ : 40 nM		+++ IC ₅₀ : 40 nM		+++ IC ₅₀ : 40 nM	+ IC ₅₀ : 300 nM					
JNJ-7706621	+++++ IC50: 9 nM	++++ IC50: 4 nM	++ IC50: 58 nM	+ IC50: 253 nM		++ ICso: 175 nM						Aurora A, Aurora B, VEGFR2
AZD5438	+++ IC50: 16 nM	+++++ IC50: 6 nM						+++ IC50: 20 nM				
MK-8776		++ IC50: 0.16 μM										Chk1,Chk2
PHA-793887	++ IC ₅₀ : 60 nM	+++++ IC ₅₀ : 8 nM		++ IC ₅₀ : 62 nM	++++ IC ₅₀ : 5 nM		+++++ IC ₅₀ : 10 nM	++ IC ₅₀ : 138 nM				GSK-3β
3S-181 HCI							+++ IC ₅₀ : 21 nM					
Palbociclib sethionate				+++++ IC50: 9 nM		+++ IC50: 15 nM						
A-674563		++ Ki: 46 nM										Akt1,PKA,GSK-3β
abemaciclib				++++ IC50: 2 nM		++++ IC50: 10 nM						
BMS-265246	++++ IC ₅₀ : 6 nM	++++ IC ₅₀ : 9 nM		++ IC ₅₀ : 230 nM								
PHA-767491	++ IC ₅₀ : 250 nM	++ IC ₅₀ : 240 nM			+ IC ₅₀ : 460 nM			+++ IC ₅₀ : 34 nM		++++ IC ₅₀ : 10 nM	+++++ IC ₅₀ : 10 nM	GSK-3β,MK2,PLK1
Milciclib	+ IC50: 398 nM	+++ IC50: 363 nM		++ ICso: 160 nM	+ IC50: 265 nM		++ ICso: 150 nM					TrkA
R547	++++ Ki: 2 nM	++++ Ki: 3 nM		++++ Ki: 1 nM								GSK-3β
NU6027	+ Ki: 2.5 μΜ	+ Κ _ί : 1.3 μΜ										ATR,DNA-PK
P276-00	++ IC ₅₀ : 79 nM	++ IC ₅₀ : 224 nM		++ IC ₅₀ : 63 nM		+ IC ₅₀ : 396 nM	+ IC ₅₀ : 2.87 μΜ	+++ IC ₅₀ : 20 nM				GSK-3β,PKCα,c-Sra
Kenpaullone	+ IC50: 0.4μM	+ IC50: 0.68µM			+ IC50: 0.85μM							GSK-3β,ERK2,c-Sr
K03861		+++++ Kd: 15.4 nM										
THZ1 2HCI							++++ IC50: 3.2 nM					
AT7519 HCI	++ IC ₅₀ : 210 nM	++ IC ₅₀ : 47 nM	+ IC ₅₀ : 360 nM	++ IC ₅₀ : 100 nM	+++ IC ₅₀ : 13 nM	++ IC ₅₀ : 170 nM	+ IC ₅₀ : 2.4 μM	++++ IC ₅₀ : <10 nM				GSK-3β
Purvalanol A		+++ IC ₅₀ : 70 nM		+ IC ₅₀ : 850 nM							++++ IC ₅₀ : 4 nM	
Ro-3306	+++ Ki: 20 nM											PKCð,SGK,ERK
SU9516	+++ IC50: 40 nM	+++ IC50: 22 nM		++ ICso: 200 nM								PDGFR
XL413										++++ IC50: 3.4 nM	++++ IC50: 3.4 nM	Pim1,CK2
LDC000067	+ IC ₅₀ : 5.513 μΜ	+ IC50: 2.441 μM		+ IC ₅₀ : 9.242 μM				+++ IC ₅₀ : 44 nM				
ML167									++ IC ₅₀ : 1522 nM			
TG003									+++ IC50: 15 nM			
Ribociclib				1								

Notes:

SAHA, 1µl

1. For more details, such as half maximal inhibitory concentrations (IC∞s) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

S1116 Palbociclib (PD-0332991) HCl

Palbociclib (PD-0332991) HCl is a highly selective inhibitor of CDK4/6 with IC50 of 11 nM/16 nM in cell-free assays, respectively. It shows no activity against CDK1/2/5, EGFR, FGFR, PDGFR, InsR, etc. Phase 3.

Size 5 mg 10 mg 50 mg



S1153 Roscovitine (Seliciclib, CYC202)

Successive States

■GAPDH

IP: cyclin E WB: Cx43

Roscovitine (Seliciclib, CYC202) is a potent and selective CDK inhibitor for Cdc2, CDK2 and CDK5 with IC50 of 0.65 μM, 0.7 μM and 0.16 μM in cell-free assays. It shows little effect on CDK4/6. Phase 2.



Circ Res, 2012, 111(2): 201-11 J Neurosci, 2012, 32(32): 11050-66

Data from [Circ Res, 2012, 111(2): 201-111 Roscovitine purchased from Selleck Cell Cycle

78

Size

HO N N N

Cell Cycle

CDK2 selecti



SNS-032 has firstly been described as a selective inhibitor of CDK2 with IC50 of 48 nM and is 10- and 20-fold selective over CDK1/CDK4. It is also found to be sensitive to CDK7/9 with IC50 of 62 nM/4 nM, with little effect on CDK6. Phase 1.



S1524 AT7519

Size

Size

S2768 Dinaciclib (SCH727965)

Dinaciclib (SCH727965) is a novel and potent CDK inhibitor for CDK2. CDK5, CDK1 and CDK9 with IC50 of 1 nM, 1 nM, 3 nM and 4 nM in cell-free assays, respectively. It also blocks thymidine (dThd) DNA incorporation. Phase 3. Size 5 mg 25 mg 50 mg 10 mM/1 mL



S1230 Flavopiridol (Alvocidib)

Flavopiridol (Alvocidib) competes with ATP to inhibit CDKs including CDK1, CDK2, CDK4 and CDK6 with IC50 of ~ 40 nM. It is 7.5-fold more selective for CDK1, 2, 4, 6 versus CDK7. Flavopiridol is initially found to inhibit EGFR and PKA. Phase 1/2.



Size 5 mg 25 mg

S7461 LDC000067 (LDC067)

LDC000067 is a highly selective CDK9 inhibitor with IC50 of 44 nM 55/125/210/ >227/ >227-fold selectivity over CDK2/1/4/6/7. J.O. M.

Size 10 mg 50 mg

S7440 Ribociclib (LEE011)

Ribociclib (LEE011) is an orally available, and highly specific CDK4/6 inhibitor. Phase 3. rober Size 5 mg 10 mg

80



JNJ-7706621 is pan-CDK inhibitor with the highest potency on CDK1/2 with IC50 of 9 nM/4 nM and shows >6-fold selectivity for CDK1/2 than for CDK3/4/6 in cell-free assays. It also potently inhibits Aurora A/B and has no activity on Plk1 and Wee1. Size

AT7519 is a multi-CDK inhibitor for CDK1, 2, 4, 6 and 9 with IC50 of

10-210 nM. It is less potent to CDK3 and little active to CDK7. Phase 2.

Flavopiridol HCl competes with ATP to inhibit CDKs including CDK1,

CDK2, CDK4 and CDK6 with IC50 of ~ 40 nM in cell-free assays. It is

7.5-fold more selective for CDK1/2/4/6 than for CDK7. Flavopiridol is

Product Citations (9):

Proc Natl Acad Sci USA, 2011, 108(20)

-

S G

5 mg 10 mg 25 mg 10 mM/1 mL

initially found to inhibit EGFR and PKA. Phase 1/2.

10 mg 25 mg 50 mg 10 mM/1 mL



S2621 AZD5438

AZD5438 (µmol/L

 \sim

NS

YBD-1e

NS

AZD5438 is a potent inhibitor of CDK1/2/9 with IC50 of 16 nM/6 nM/20 nM in cell-free assays. It is less potent to CDK5/6 and also inhibits GSK3_β. Phase 1. Size 10 mg 50 mg 10 mM/1 mL CDK9 selective





S2735 MK-8776 (SCH 900776) Chk1 selectiv

MK-8776 (SCH 900776) is a selective Chk1 inhibitor with IC50 of 3 nM in a cell-free assay. It shows 500-fold selectivity against Chk2. Phase 2. ----- Page 81



BS-181 HCl is a highly selective CDK7 inhibitor with IC50 of 21 nM. It is more than 40-fold selective for CDK7 than for CDK1, 2, 4, 5, 6, or 9.



Data from [Arthritis Rheumatol, 2014, 66(6): 1537-46] BS-181 HCI purchased from Selleck

S1579 Palbociclib (PD0332991) Isethionate

Palbociclib (PD0332991) Isethionate is a highly selective inhibitor of CDK4/6 with IC50 of 11 nM/16 nM in cell-free assays. It shows no activity against CDK1/2/5, EGFR, FGFR, PDGFR, InsR, etc. Phase 3. Size 10 mg 25 mg 50 mg



S7158 abemaciclib (LY2835219)

LY2835219 is a potent and selective inhibitor of CDK4 and CDK6 with



S2742 PHA-767491 (CAY10572)

PHA-767491 is a potent ATP-competitive dual Cdc7/CDK9 inhibitor with IC50 of 10 nM and 34 nM in cell-free assays, respectively. It displays ~20-fold selectivity against CDK1/2 and GSK3-β, 50-fold selectivity against MK2 and CDK5, 100-fold selectivity against PLK1 and CHK2. Size 10 mg 50 mg 10 mM/1 mL



S1532 AZD7762



Data from [Cancer Discov, 2012, 2(6)

AZD7762 purchased from Selleck

Product Citations (16)

Nat Biotechnol, 2011, 29(6): 542-6

Cancer Discov, 2012, 2(6): 524-39

MK-8776 (SCH 900776) is a selective Chk1 inhibitor with IC50 of 3 nM in a cell-free assay. It shows 500-fold selectivity against Chk2. Phase 2. Size



524-391

Chk Inhibitors

S2670 A-674563

S7747 Ro-3306

S7549 THZ1 2HCI

selectivity for CDK7.

5 mg 25 mg

10 mg 50 mg 200 mg

Size

Size

Inhibitory Selectivity

Inhibitor Name	Chk1	Chk2	Other
AZD7762	+++ IC50: 5 nM		
LY2603618	+++ IC50: 7 nM		
MK-8776	+++ IC50: 3 nM	+ ICso: 1.5 μM	CDK2
CHIR-124	++++ IC50: 0.3 nM	+ IC50: 697.4 nM	FLT3,PDGFR,GSK-3
PF-477736	++++ Ki: 0.49 nM	++ Ki: 47 nM	VEGFR2,Fms,YES
SAR-020106	++ IC50: 13.3 nM		
Prexasertib	++++ Ki: 0.9 nM	++ IC50: 8 nM	RSK

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

AZD7762 is a potent and selective inhibitor of Chk1 with IC50 of 5 nM in

a cell-free assay. It is equally potent against Chk2 and less potent

against CAM, Yes, Fyn, Lyn, Hck and Lck. Phase 1.

▲6-Tubulin

<β-Tubulin

Size 2 mg 25 mg 100 mg 10 mM/1 ml

- - + + + + - + AZD7762 - + + - - + + MK-1775

---- Cyclin B1

---- <--- <CDK1 total

0.7 0.5 0.3 0.2 1.0 1.0 1.0 0.5 Ratio pY15/total CDK

---- 4CDK1 total

0.7 0.6 0.2 0.1 0.6 0.5 0.1 0.0 Ratio pY15/total CDK1

S2735 MK-8776 (SCH 900776)

---- <Cyclin B1

--- - <-- <CDK1 pY15

A-674563 is an Akt1 inhibitor with Ki of 11 nM in cell-free assays,

RO-3306 is an ATP-competitive, and selective CDK1 inhibitor with Ki of

THZ1 is a covalent CDK7 inhibitor which has the unprecedented ability to target a remote cysteine residue located outside of the canonical

kinase domain, providing an unanticipated means of achieving

20 nM, >15-fold selectivity against a diverse panel of human kinases.

----- Page 13

modest potent to PKA and >30-fold selective for Akt1 over PKC.



Product Citations (10) Proc Natl Acad Sci USA, 2013, 110(10): Nucleic Acids Res, 2013, 41(22):

Data from [Nucleic Acids Res, 2013, 41(22): 10334-44] PD0332991 purchased from Selleck

IC50 of 2 nM and 10 nM in cell-free assays, respectively. Phase 3. Size 5 mg ----

Biochem J, 2014, 459(3): 513-24 Data from [Biochem J, 2014, 459(3): LY2835219 (219) purchased from

d a

Chk / ROCK

S2626 LY2603618 (IC-83)

LY2603618 is a highly selective Chk1 inhibitor with potential anti-tumor activity in a cell-free assay. IC₅₀=7 nM, showing approximately 100-fold more potent against Chk1 than against any of the other protein kinases evaluated.

Size 5 mg 10 mg 50 mg 10 mM/1 mL

U937 (8h)	
Craft ^d - 5 th c ⁴ c ⁴ b ⁴ c	Product Citations (9): J Pathol, 2015, 10.100 Cancer Lett, 2014, 356
1.00 1.29 2.35 5.07 ← CDK1 ← β-actin	Data from [J Hematol 7(1): 53] LY2603618 purchased

Y-39 Neta GSK Ripa hvd 02/path.4528 dihy 6(2 Pt B): 656-68 KD0 AT1 I Oncol. 2014

Chk1 sele

Chk1 selective

Chk1 selective

from Selleck

S2683 CHIR-124

CHIR-124 is a novel and potent Chk1 inhibitor with IC50 of 0.3 nM in a cell-free assay. It shows 2,000-fold selectivity against Chk2, 500- to 5,000-fold less activity against CDK2/4 and Cdc2. $\operatorname{Cr}^{n,N}_{\mathcal{A}}$



667-781 CHIR-124 purchased from Selleck

S2904 PF-477736 (PF-736, PF-00477736)

PF-477736 is a selective, potent and ATP-competitive Chk1 inhibitor with K of 0.49 nM in a cell-free assay and also inhibits VEGFR2, Aurora-A, FGFR3, Flt3, Fms (CSF1R), Ret and Yes. It shows ~100-fold selectivity for Chk1 than Chk2. Phase 1.



S7178 Prexasertib (LY2606368)

Prexasertib (LY2606368) is an ATP-competitive CHK1 inhibitor with a Ki value of 0.9 nmol/L. For CHK2 and RSK, its IC50 values are 8 nM and 9 nM respectively in cell-free assay

Size 2 mg 5 mg 25 mg

ROCK Inhibitors

Inhibitory Selectivity

Inhibitor Name	ROCK	ROCK1	ROCK2	Other	
Y-27632 2HCI		+ Ki: 140 nM	+ Ki: 300 nM		
Thiazovivin	+ IC50: ~0.5 μM				
Fasudil HCI			+ K _i : 330 nM	PKA,PKG,PKC	
GSK429286A		+++ IC50: 14 nM	++ IC50: 63 nM		

Inhibitory Selectivity

Inhibitor Name	ROCK	ROCK1	ROCK2	Other
RKI-1447		+++ IC50: 14.5 nM	+++ IC50: 6.2 nM	
Y-39983 HCI	++++ K _i : 2 nM			
Netarsudil 2HCI				norepinephrine transporter (NET)
GSK269962 HCI		++++IC50: 1.6 nM	++++IC50: 4 nM	MSK1,RSK1
Ripasudil hydrochloride dihydrate		++ IC ₅₀ : 51 nM	+++ IC ₅₀ : 19 nM	
KD025			++ IC50: 60 nM	
AT13148		+++ IC50: 6 nM	++++IC50: 4 nM	PKA,p70S6K,Akt1

Notes: 1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

S1049 Y-27632 2HCI

Size

Y-27632 2HCl is a selective ROCK1 (p160ROCK) inhibitor with Ki of 140 nM in a cell-free assay, exhibiting >200-fold selectivity over other kinases, including PKC, cAMP-dependent protein kinase, MLCK and PAK.



Nature, 2016, 532(7597): 107-11 Cell Res. 2013. 23(10): 1187-200 Data from I J Control Release, 2013

Y-27632 2HCI purchased from Selleci

S1459 Thiazovivin

ETer 150

100

20.6

enea

음0.2

Thiazovivin is a novel ROCK inhibitor with IC50 of 0.5 µM in a cell-free assay, promoting hESC survival after single-cell dissociation. Size



S1573 Fasudil (HA-1077) HCI ROCK2 selective Fasudil (HA-1077), a potent and selective inhibitor of Rho kinase, displays less potent inhibiton over PKA, PKG, PKC and MLCK with K of 1.6, 1.6, 3.3, and 36 µM in cell-free assays, respectively.

Size 200 mg 500 mg 10 mM/1 mL 4T1 Product Citations (7): Biosens Bioelectron, 2016, 86: 508-15 PBS Fasudi J Clin Invest, 2014, 124(4): 1646-59 Data from [J Clin Invest, 2014, 124(4);

1646-591 Fasudil HCI purchased from Selleck

12m



83

Wee1 / Rho / c-Myc / PD-1/PD-L1



MK-1775 pt

ne

S8148 PD0166285

PD0166285 is a potent Wee1 and Chk1 inh nanomolar concentrations.PD0166285 is a r abrogator.

Cell Cycle Size 5 mg 25 mg

		0.13,~20
	S7686 ML141	
Mol Cancer Ther, 2012, 22] urchased from Selleck	ML141 (CID-2950007), is demonstrated to be a poreversible non-competitive inhibitor of Cdc42 GTP vitro assays, with ICs0 of 200 nM and selectivity agai of the Rho family of GTPases (Rac1, Rab2, Rab7).	tent, selective and ase suitable for in inst other members
nibitor with activity at novel G2 checkpoint	Size 5 mg 25 mg 100 mg	""Far
ptoto.	S7719 CCG-1423	<i>w</i>
	CCG-1423 is a specific RhoA pathway inhibitor, w mediated transcription.	hich inhibits SRF-
	Size 10 mg 50 mg 200 mg	- Artha

S7319 EHop-016

Rho Inhibitors

Inhibitory Selectivity

Inhibitor Name	Rho	c-myc ini
EHT 1864	+++ K _d : 50 nM	S7153 10058-F4
Zoledronic Acid	4	10058-F4 is a c-Myc
K-Ras(G12C) inhibitor 9	4	expression.
K-Ras(G12C) inhibitor 6	4	Size 25 mg
K-Ras(G12C) inhibitor 12	√	
6H05	V	

Notes:

1. For more details, such as half maximal inhibitory concentrations (IC $_{\rm 50}{\rm s})$ and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation. 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

S1314 Zoledronic Acid (Zoledronate, CGP-4244)

Zoledronic acid (ZA), a potent osteoclast inhibitor, induces apoptosis in osteoclasts by inhibiting enzymes of the mevalonate pathway and preventing the isoprenylation of small GTP-binding proteins such as Ras and Rho

Size 25 mg 100 mg

S8031 NSC 23766

Size 5 mg 25 mg

84

NSC 23766 is an inhibitor of Rac GTPase targeting Rac activation by guanine nucleotide exchange factors (GEFs) with IC $_{50}$ of ~50 μM in a cell-free assay; does not inhibit the closely related targets. Cdc42 or RhoA. Size 10 mg 50 mg 10 mM/1 mL

S7331 K-Ras(G12C) inhibitor 12

K-Ras(G12C) inhibitor 12 is an allosteric inhibitor of oncogenic K-Ras(G12C). ${}^{\rm a}_{\rm p} = {}^{\rm a}_{\rm p} = {}^{$

hibitor inhibitor that specifically inhibits the c-Myc-Max events transactivation of c-Myc target gene D The

PD-1/PD-L1 Inhibitors

S7912 PD-1/PD-L1 inhibitor 2

PD-1/PD-L1 inhibitor 2 is a small-molecule PD-1/PD-L1 interaction inhibitor with IC50 of 18 nM.

appine.

S7911 PD-1/PD-L1 inhibitor 1

Size 5 mg 25 mg

PD-1/PD-L1 inhibitor 1 is a small-molecule inhibitor of PD-1/PD-L1 interaction with IC₅₀ of 6 nM. Size 5 mg 25 mg appiro

S8158 PD-1/PD-L1 Inhibitor 3

PD-1/PD-L1 Inhibitor 3 (Programmed Death-1/Programmed Death -Ligand 1 Inhibitor 3) is a Macrocyclic inhibitor of PD-1/PD-L1 ΩľΎ interaction with IC50 of 5.6 nM. Size 1 mg 5 mg



mitab

700

TGF-beta/Smad Pathway



Bcr-Abl Inhibitors

Detailed product information is on page 51-52

PKC Inhibitors

Detailed product information is on page 72-73

ROCK Inhibitors

Detailed product information is on page 82-83

TGF-beta/Smad Inhibitors

Inhibitory Selectivity

Inhibitor Name ALK1 ALK3 TGF^βRI/ALK5 TGFBRII TGF-B Smad3 Other ALK2 ALK4 ALK6 SB431542 ++ IC50: 94 nM I DN-193189 ++++ ICso: 5 nM ++ ICso: 30 nM Galunisertib ++ IC50: 56 nM LY2109761 +++ Ki: 38 nM K_i: 300 nM SB525334 +++ IC50: 14.3 nM SB505124 ++ IC50; 47 nM IC50: 129 nM GW788388 +++ IC50: 18 nM RIPK2,CK1ō, LY364947 IC50: 59 nM IC50: 0.4 µM MLK-7K RepSox ++++ IC50: 4 nM LDN-193189 HC ++++ ICso: 5 nM +++ ICso: 30 nM K02288 ++++ IC50: 1.8 nM +++++ IC50: 1.1 nM +++ IC50: 34.4 nM + IC50: 302 nM IC50: 321 nM ++++IC50: 6.4 nN LDN-214117 +++ IC50: 24 nM SD-208 ICso: 48 nM EW-7197 + IC50: 13 nM ++++ IC50: 11 nM ML347 IC50: 46 nM +++ IC50: 32 nM IC50: 10.8 µM IC₀₀: 9.83 µM I DN-212854 +++ IC50: 2.4 nM ++++ IC50: 1.3 nM IC50: 85.8 nM IC50: 2133 nM + IC50: 9276 nM DMH1 ++ ICso: 107.9 nM

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TGF-beta/Smad

Inhibitory Selectivity

Inhibitor Name	ALK1	ALK2	ALK3	ALK4	TGFβRI/ALK5	ALK6	TGFβRII	TGF-β	Smad3	Other
Pirfenidone								1		
SIS3 HCI									1	
Hesperetin								1		Histamine receptor
Notes:										

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

S1067 SB431542

pSMAD3

SMAD3

86

GF-beta/Smad

S2704 LY2109761 TGF\$RI/ALK5 selective

SB431542 is a potent and selective inhibitor of ALK5 with IC50 of 94 nM in a cell-free assay, 100-fold more selective for ALK5 than for p38 MAPK and other kinases.

5 mg 10 mg 10 mM/1 mL Size 10 mg 50 mg 10 mM/1 mL Size S QQ__C AP-O-30 Product Citations (6) 25 Connect Tissue Res. 2015. 56(4): 20 288-99 15 Toxicology, 2014, 326C: 9-17 Product Citations (24): 10 nature, 2016, 10, 1038/nature17408 days of treatment (TGFß1 10ng/ml) J Clin Invest, 2015, 125(2): 796-808 5 CD34⁺ HSPCs Data from [Toxicology, 2014, 326C; galangin (148mM) 9-171 - miR-99 LY2109761 LY2109761 purchased from Selleck miR-let7c - miR-125b ** - tricistron S1476 SB525334 TGF\$RI/ALK5 selective -@ - SB431542 Data from [Genes Dev. 2014, 28(8); SB525334 is a potent and selective inhibitor of TGF^β receptor I (ALK5) 2 4 6 8 10 -9- sh-SMAD4 858-741 days of treatment (TGFß1 1ng/ml) with IC50 of 14.3 nM in a cell-free assay, is 4-fold less potent to ALK4 SB431542 purchased from Selleck than ALK5 and inactive to ALK2, 3, and 6. 5 mg 50 mg 100 mg 10 mM/1 mL S2618 LDN-193189 (DM3189) Size LDN-193189 is a selective BMP signaling inhibitor, inhibiting the 0.3% NaCl 8.0% NaC transcriptional activity of the BMP type I receptors ALK2 and ALK3 with 334 Vehicle SB525334 (10 mg/kg/d ehicle SB5 IC50 of 5 nM and 30 nM in C2C12 cells, respectively, exhibiting 200-fold ----selectivity for BMP versus TGF-β. Smad2/3 Product Citations (7): Size 2 mg 5 mg 25 mg Cancer Lett. 2014. 355(1): 130-40 PTEN Hypertension, 2013, 62(5); 951-6 WT MUT LDN193189 LDN193189 Product Citations (12): Ak J Clin Invest. 2015, 125(2): 796-808 - 100nM 200nM - 100nM 200nM p-NOS3 (S1177) Cancer Cell, 2014, 26(4): 521-33 Data from [Hypertension, 2013, 62(5): ----nad1/5 951-61 GAPDH SB525334 purchased from Selleck -tubulir ----Data from [J Cell Sci, 2012, 126 (Pt 1): 234-431 S7146 DMH1 ALK2 selective LDN-193189 purchased from Selleck DMH1 is a selective BMP receptor inhibitor with IC50 of 107.9 nM for ALK2, exhibiting no inhibition on AMPK, ALK5, KDR (VEGFR-2) or S2230 Galunisertib (LY2157299) TGFβRI/ALK5 selective PDGFR. Galunisertib (LY2157299) is a potent TGF_β receptor I (T_βRI) inhibitor Size 10 mg 25 mg with IC50 of 56 nM in a cell-free assay. Phase 2/3. Size 5 mg 10 mg 50 mg 10 mM/1 mL ŭ U Him HLE cells S7507 LDN-193189 HCI TGFβR1/2 inhibitor (LY2157299) DMSO 100 nmol/L 500 nmol/L LDN193189 HCI is the hydrochloride salt of LDN193189, which is a GLI2 selective BMP signaling inhibitor, and inhibits the transcriptional activity of the BMP type I receptors ALK2 and ALK3 with IC50 of 5 nM and 30 nM nSMAD3 Product Citations (5): in C2C12 cell lines, respectively, 200-fold selectivity for BMP versus Sci Rep, 2016, 6:23056 SMAD3 TGF-β. Cancer Res. 2014. 74(21): 5963-77 HLF cells TGF8R1/2 inhibito Size 5 mg 10 mg 50 mg (LY2157299) DMSO 100 nmol/L 500 nmol/L GLI2

> Data from [Cancer Res 2014 10.1158/0008-5472.CAN-14-02251 LY2157299 purchased from Sellect



LY2109761 is a novel selective TGF-ß receptor type I/II (TßRI/II) dual

inhibitor with K of 38 nM and 300 nM in cell-free assay, respectively;

shown to negatively affect the phosphorylation of Smad2.

Size 10 mg 50 mg 10 mM/1 mL

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S2907 Pirfenidone (S-7701, AMR-69)

TGF-B selective

Pirfenidone is an inhibitor for TGF- β production and TGF- β stimulated collagen production, reduces production of TNF- α and IL-1 β , and also has anti-fibrotic and anti-inflammatory properties. Phase 3. Size 10 mg 50 mg 10 mM/1 mL

S2805 LY364947

LY364947 is a potent ATP-competitive inhibitor of TGFBR-I with IC50 of 59 nM in a cell-free assay, showing 7-fold selectivity over TGF β R-II. Size 10 mg 25 mg 50 mg



signaling by suppressing Smad3 phosphorylation without affecting the MAPK/p38, ERK, or PI3-kinase signaling pathways.

Size

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TGF-beta/Smad

TGFBRI/ALK5 selective

50



S7223 RepSox (E-616452, SJN 2511)

RepSox is a potent and selective inhibitor of the TGFBR-1/ALK5 with IC50 of 23 nM and 4 nM for ATP binding to ALK5 and ALK5 autophosphorylation in cell-free assays, respectively.

Size 10 mg 25 mg 0-0

S7959 SIS3 HCI

SIS3, a novel specific inhibitor of Smad3, inhibits TGF-B and activin

2 mg 5 mg 25 mg

Product Citation (1): Chem Biol Interact, 2014, 217: 1-8 Data from [Chem Biol Interact, 2014,

DNA Damage



HDAC Inhibitors

Detailed product information is on page 19-23

Detailed product information is on page 29-30

Sirtuin Inhibitors | Activators

ATM/ATR Inhibitors | Activator

Detailed product information is on page 15-16

Detailed product information is on page 18

DNA-PK Inhibitors

PARP Inhibitors

Detailed product information is on page 23-24

DNA/RNA Synthesis Inhibitors | Antagonist | Chemical | Modulator

DNA/RNA Synthesis Inhibitors

S1166 Cisplatin

Cisplatin is an inorganic platinum complex, which is able to inhibit DNA synthesis by conforming DNA adducts in tumor cells. Size 50 mg NHa

S1149 Gemcitabine HCI Gemcitabine HCI is a DNA synthesis inhibitor with IC50 of 50 nM, 40 nM, 18 nM and 12 nM in PANC1, MIAPaCa2, BxPC3 and Capan2 cells, respectively



S2684 CX-5461

Bleomycin Sulfate is a glycopeptide antibiotic and an anticancer agent for squamous cell carcinomas (SCC) with IC50 of 4 nM in UT-SCC-19A cells

Selleck

Product Citations (6): Cancer Cell, 2013, 24(5): 617-30

1839-44

20(14): 3849-611 Carboplatin purchased from Selleck

Product Citations (6):

Int J Cancer, 2014, 136(4): E51-61

Data from [Int J Cancer, 2014

Oxaliplatin purchased from Selleck

10.1002/ijc.29161]

Proc Natl Acad Sci USA, 2015, 112(6);

Data from [Clin Cancer Res, 2014,

Carboplatin is a DNA synthesis inhibitor by binding to DNA and

interfering with cell repair mechanism in A2780, SKOV-3, IGROV-1, and

Oxaliplatin inhibits DNA synthesis by conforming DNA adducts in RT4,

TCCSUP, A2780, HT-29, U-373MG, U-87MG, SK-MEL-2, and HT-144

0.5 1.3 2.5 5 10 20

Sofosbuvir (PSI-7977, GS-7977) is a HCV NS5B polymerase inhibitor

Pemetrexed is a novel antifolate and antimetabolite for TS, DHFR and

Fludarabine is a STAT1 activation inhibitor which causes a specific

depletion of STAT1 protein (and mRNA) but not of other STATs. Also a

Capecitabine is a tumor-selective fluoropyrimidine carbamate which

achieves higher intratumoral 5-FU level with lower toxicity than 5-FU.

for the treatment of chronic hepatitis C virus (HCV) infection.

GARFT with K of 1.3 nM, 7.2 nM and 65 nM, respectively.

DNA synthesis inhibitor in vascular smooth muscle cells.

6,2 12,5 25 50 100 Etoposida (utili)



S1215 Carboplatin (JM-8, CBDCA, NSC 241240)

Size 50 mg 100 mg 200 mg

S1224 Oxaliplatin (L-OHP)

50 mg 100 mg 200 mg

S2794 Sofosbuvir (PSI-7977, GS-7977)

Size 5 mg 25 mg 100 mg

S1135 Pemetrexed (LY-231514)

S1156 Capecitabine

S1491 Fludarabine (FaraA, Fludarabinum)

Size 50 mg 200 mg 1 g 10 mM/1 mL

HX62 cells.

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cells.

Size

S1214 Bleomycin Sulfate (NSC125066)

CX-5461 is an inhibitor of rRNA synthesis, selectively inhibits Pol I-driven transcription of rRNA with IC50 of 142 nM in HCT-116, A375, and MIA PaCa-2 cells, has no effect on Pol II, and possesses 250- to 300-fold selectivity for inhibition of rRNA transcription versus DNA replication and protein translation.

Size 5 mg 10 mg 50 mg



S1209 Fluorouracil (5-Fluoracil, 5-FU, NSC 19893)

Fluorouracil (5-Fluoracil, 5-FU) is an DNA/RNA synthesis inhibitor, which interrupts nucleotide synthetic by inhibiting thymidylate synthase (TS) in tumor cells.

Size 100 mg 200 mg 10 mM/1 mL

S1648 Cytarabine

Cytarabine (Cytosine arabinoside, AraC) is an antimetabolic agent and DNA synthesis inhibitor with IC50 of 16 nM in wild-type CCRF-CEM cells.

Size 50 mg 5 g

S1714 Gemcitabine

Gemcitabine, a nucleic acid synthesis inhibitor, is a very potent and specific deoxycytidine analogue, used as chemotherapy. Size 50 mg 10 mM/1 mL



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Product Citations (13) Sci Transl Med, 2015, 7(284): 284ra57 Proc Natl Acad Sci USA, 2015, 112(6): 1830-44

42(10): 6436-47] Gemcitabine purchased from Selleck

S1218 Clofarabine

Clofarabine inhibits the enzymatic activities of ribonucleotide reductase (IC50 = 65 nM) and DNA polymerase. .XX

Size 10 mg 50 mg 10 mM/1 mL

S1192 Raltitrexed (ZD-1694)

Raltitrexed is a thymidylate synthase inhibitor with an IC50 of 9 nM for the inhibition of L1210 cell growth.

Size 10 mg 50 mg 100 mg 10 mM/1 mL

S1302 Ifosfamide (NSC109724, Isophosphamide)

Ifosfamide is a nitrogen mustard alkylating agent used in the treatment of cancer.

Size Juliy Tulliv/Thic

S7742 SCR7 SCR7 is a specific DNA Ligase IV inhibitor, which blocks nonhomologous end-joining

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S1221 Dacarbazine (DTIC-Dome)

Dacarbazine is a triazene derivative with antineoplastic activity. Dacarbazine alkylates and cross-links DNA during all phases of the cell cycle, resulting in disruption of DNA function, cell cycle arrest, and apoptosis; used in the treatment of various cancers.

Size 50 mg 10 mM/1 mL

S7419 Blasticidin S HCI

Blasticidin S HCl is a nucleoside antibiotic isolated from Stretomyces girseochromogenes, and acts as a DNA and protein synthesis inhibitor, used to select transfected cells carrying bsr or BSD resistance genes.

Size 25 mg 100 mg

istance genes.	DNA	/RNA	Sy	nthes	sis Cł	nem	ic	al
in the second second	S1982	Adenine su	Ilfate					
на	Adenine	sulfate is a	sulfate	salt form	of adenine	which	is a	purine

N NN-N

S2504 Ribavirin

Ribavirin, a synthetic guanosine analogue, possesses a broad spectrum of activity against DNA and RNA viruses. Size 100 mg 200 mg 10 mM/1 mL HO YON

S8146 Mitomycin C

Mitomycin C is an antineoplastic antibiotic by inhibiting DNA synthesis, used to treat different cancers

Size 10 mg 50 mg 200 mg



derivative and a nucleobase with a variety of roles in biochemistry.

DNA/RNA Synthesis Antagonist

Flupirtine maleate is the salt form of Flupirtine, which is a centrally

acting non-opioid analgesia, is a selective neuronal potassium channel

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H₂SO₄

opener that also has NMDA receptor antagonist properties.

Size 10 mg 25 mg 100 mg 10 mM/1 mL

S1334 Flupirtine maleate

Size 50 mg 5 g

Topoisomerase Inhibitors

Inhibitory Selectivity

Inhibitor Name	Topoisomerase	Торо І	Topo II	Topo IV	Other
Camptothecin		++ IC ₅₀ : 0.68 μM			
Topotecan HCI		++++ IC50: 13 nM			
Idarubicin HCI			+++ IC50: 3.3 ng/mL		Multicellular spheroids
Daunorubicin HCI	+++ K: 20 nM				
Betulinic acid		++ IC ₅₀ : 5 μM			HIV-1, Aminopeptidase N
Flumequine			+ IC ₅₀ : 15 μΜ		
Doxorubicin			1		
Etoposide			1		
Irinotecan		V			
Epirubicin HCI	1				
Mitoxantrone HCI			1		
Moxifloxacin HCI			√		
Irinotecan HCI Trihydrate		V			
SN-38		V			
Amonafide			1		
Teniposide			√		
Gatifloxacin	\checkmark				
Genistein			1		EGFR
Mitoxantrone			1		
Levofloxacin			1		
Pirarubicin			1		
Ciprofloxacin				1	

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.



Topoisomerase / Telomerase / DNA Alkylator S4908 SN-38



Size 5 mg 10 mg 10 mM/1 mL

Melphalan is a phenylalanine derivative of nitrogen mustard with antineoplastic activity Size 100 mg 500 mg

Stem Cells and Wnt Pathway



GSK-3 Inhibitors

Detailed product information is on page 14-15

JAK Inhibitors

Detailed product information is on page 24-26

STAT Inhibitors

Detailed product information is on page 63-64



Inhibitory Selectivity

Inhibitor Name	γ secretase	Αβ	Notch	Other Targets
Dibenzazepine	+++ IC ₅₀ : 2.6 nM		+++ IC50: 2.9 nM	
LY411575	++++ IC50: 0.082 nM		++++ IC ₅₀ : 0.39 nM	
L-685,458	+ Ki: 17 nM			
FLI-06			+ EC50: 2.3 μM	
LY3039478			++++ IC50: ~1 nM	
PF-03084014	++ IC50: 6.2 nM			
MK-0752		V		Αβ

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2 "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

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AB selective

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

S2215 DAPT (GSI-IX)

DAPT (GSI-IX) is a novel $\gamma\text{-secretase}$ inhibitor, which inhibits $A\beta$ production with IC50 of 20 nM in HEK 293 cells.





S1575 RO4929097

RO4929097 is a y secretase inhibitor with IC50 of 4 nM in a cell-free assay, inhibiting cellular processing of A β 40 and Notch with EC₅₀ of 14 nM and 5 nM, respectively. Phase 2.

Semagacestat (LY450139) is a v-secretase blocker for AB42, AB40 and

A\beta38 with IC $_{50}$ of 10.9 nM, 12.1 nM and 12.0 nM, also inhibits Notch

signaling with IC50 of 14.1 nM in H4 human glioma cell. Phase 3.

50 mg 10 mM/1 mL

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Αβ 1-42 Αβ x-42

Dibenzazepine (YO-01027) is a dipeptidic γ -secretaseinhibitor with IC $_{50}$

of 2.6 nM and 2.9 nM in cell-free assays for APPL and Notch cleavage,





S1594 Semagacestat (LY450139)

10 mg 5 mg

Semagacestat (1µM)

Aß detection

respectively

Substrate accumulation

S2711 Dibenzazepine (YO-01027)

Size 2 mg 5 mg 25 mg 10 mM/1 mL

Data from [Dev Cell, 2014, 30(4): 410-221 RO4929097 (RO) purchased from

Product Citations (3):

1540-50]

J Biol Chem. 2014, 289(3): 1540-50 J Biol Chem, 2012, 287(15): 11810-9

Data from [J Biol Chem, 2014, 289(3):

Semagacestat purchased from Selleck

Avagacestat (BMS-708163) is a potent, selective, orally bioavailable γ-secretase inhibitor of Aβ40 and Aβ42 with IC50 of 0.3 nM and 0.27 nM, demonstrating a 193-fold selectivity against Notch. Phase 2.



S2714 LY411575

Size

LY411575 is a potent y-secretase inhibitor with IC50 of 0.078 nM/0.082 nM (membrane/cell-based), also inhibits Notch clevage with IC50 of 0.39 nM in APP or N∆E expressing HEK293 cells.

5 mg 10 mg 50 mg 10 mM/1 mL Size

S1262 Avagacestat (BMS-708163)



LY3039478 is an oral Notch inhibitor with an IC50 of 0.41 nM. 5 mg 25 mg

Hedgehog/Smoothened Inhibitors Agonists | Antagonists

Inhibitory Selectivity

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Inhibitor Name	Hedgehog	Smoothened		GLI
Vismodegib	+++ IC50: 3 nM			
Cyclopamine		++ IC50: 46 nM		
Erismodegib		++++ IC ₅₀ : 1.3 nM		
PF-5274857		+++ IC50: 5.8 nM		
GANT61			+	IC50: 5 µM
SANT-1		++++ Kd: 1.2 nM		
Taladegib		V		
BMS-833923		1		
Jervine	1			

Notes: 1 For more details such as half maximal inhibitory concentrations (ICros) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

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AB selective

Gamma-secret	tase Inl	hibitors

Inhibitory Selectivity

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Inhibitor Name	γ secretase	Αβ	Notch	Other
DAPT (GSI-IX)		+ IC50: 20 nM		Αβ
RO4929097	+++ IC50: 4 nM		+++ IC50: 5 nM	Αβ40
Semagacestat		++ IC ₅₀ : 10.9 nM	++ IC ₅₀ : 14.1 nM	
Avagacestat		++++ IC50: 0.3 nM		

Cells and Wnt

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Hedgehog/Smoothened Inhibitors Casein Kinase Inhibitors

S1082 Vismodegib (GDC-0449)

Size

Vismodegib (GDC-0449) is a potent, novel and specific hedgehog inhibitor with IC50 of 3 nM and also inhibits P-gp with IC50 of 3.0 µM in a cell-free assay. Č^QiČ, 5 mg 50 mg 200 mg 10 mM/1 mL



S8075 GANT61 (NSC 136476)

GANT61 is an inhibitor for GLI1 as well as GLI2-induced transcription, inhibits hedgehog with IC50 of 5 µM in GLI1 expressing HEK293T cell, displays selectivity over other pathways, such as TNF and glucocorticoid receptor gene transactivation.

Size 10 mg 50 mg

Hedgehog/Smoothened Agonists

S3042 Purmorphamine

Purmorphamine, which directly binds and activates Smoothened, blocks BODIPY-cyclopamine binding to Smo with IC50 of ~ 1.5 μ M in HEK293T cell and also is an inducer of osteoblast differentiation with EC₅₀ of 1 µM.).).

Size 5 mg 25 mg

S7779 Smoothened Agonist (SAG) HCI

Smoothened Agonist (SAG) HCl is a cell-permeable Smoothened (Smo) agonist with EC50 of 3 nM in Shh-LIGHT2 cells.

Size 2 mg 5 mg 25 mg

Hedgehog/Smoothened Antagonists

S1146 Cyclopamine

Cyclopamine is a specific Hedgehog (Hh) signaling pathway antagonist of Smoothened (Smo) with IC50 of 46 nM in TM3Hh12 cells. Size





Inhibitory Selectivity

Inhibitor Name	CK1	CK2	Other Targets
Silmitasertib		+++ IC50: 1 nM	
D 4476	++ IC50: 300 nM		ALK5
Notes:			

1 For more details such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

S2248 Silmitasertib (CX-4945)

GLI selective Silmitasertib (CX-4945) is a potent and selective inhibitor of CK2 (casein kinase 2) with IC50 of 1 nM in a cell-free assay, less potent to Flt3, Pim1 and CDK1 (inactive in cell-based assay). Phase 1/2.



S7642 D 4476

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D 4476 is a potent, selective, and cell-permeant CK1 (casein kinase 1) inhibitor with IC50 of 200 nM and 300 nM in a cell-free assay for CK1 from Schizosaccharomyces pombe and CK1 $\!\delta\!$, respectively. Also acts as an ALK5 inhibitor with IC50 of 500 nM. Size 10 mg 50 mg 200 mg

Hippo Pathway Inhibitors

S8334 XMU-MP-1 XMU-MP-1 is an inhibitor of MST1/2 with IC50 values of 71.1±12.9 nM and 38.1±6.9 nM against MST1 and MST2, respectively. Size 2 mg 5 mg 25 mg 1XQ3 Ó.

S8164 YAP-TEAD Inhibitor 1 (Peptide 17) new

Peptide 17 is a inhibitor of this YAP-TEAD protein-protein interaction which has potential usage in treatment of YAP-involved cancers with IC50 of 25nM Size 1 mg



Ubiquitin Pathway



Proteasome Inhibitors

Inhibitory Selectivity

Inhibitor Name	Proteasome	20S proteasome
Bortezomib (PS-341)		++++ Ki: 0.6 nM
MG-132		+ ICso: 100 nM
Carfilzomib (PR-171)	+++ IC50: 5 nM	
MLN9708		+++ Ki: 0.93 nM
Ixazomib (MLN2238)		++++ K _i : 0.93 nM
ONX-0914 (PR-957)		++ ICso: ~10 nM
Oprozomib (ONX 0912)		++ ICso: 36 nM
Delanzomib (CEP-18770)		+++ IC ₅₀ : 3.8 nM
Celastrol		+ IC ₅₀ : 2.5 μM
VR23	++++ ICso: 1 nM	
PI-1840		++ ICso: 27 nM
Epoxomicin		1

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

S1013 Bortezomib (PS-341)

Bortezomib (PS-341) is a potent 20S proteasome inhibitor with K of 0.6 nM. It exhibits favorable selectivity towards tumor cells over normal cells.



S2619 MG-132

MG-132 is an inhibitor of proteasome with IC50 of 100 nM in a cell-free assay, and also inhibits calpain with IC50 of 1.2 µM. Size



siRNA

Control

www.selleckchem.com

MG-132 purchased from Selleck

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Proteasome / DUB

S2853 Carfilzomib (PR-171)

Carfilzomib (PR-171) is an irreversible proteasome inhibitor with IC $_{50}$ of <5 nM in ANBL-6 cells, displayed preferential in vitro inhibitory potency against the ChT-L activity in the β 5 subunit, but little or no effect on PGPH and T-L activities.



S2181 MLN9708

MLN9708 immediately hydrolyzed to MLN2238, the biologically ac form, on exposure to aqueous solutions or plasma. MLN2238 inhi the chymotrypsin-like proteolytic (β5) site of the 20S proteasome IC_{50}/K_i of 3.4 nM/0.93 nM in cell-free assays, less potent to β 1 and activity to $\beta 2$. Phase 3.



S2180 Ixazomib (MLN2238)

Ixazomib (MLN2238) inhibits the chymotrypsin-like proteolytic (β5) site of the 20S proteasome with IC_{50} and K_i of 3.4 nM and 0.93 nM in cell-free assays, respectively, also inhibits the caspase-like (B1) and trypsin-like (β2) proteolytic sites, with IC50 of 31 and 3500 nM. Phase 3. Size 5 mg 10 mg 10 mM/1 mL



S7172 ONX-0914 (PR-957)

ONX-0914 (PR-957) is a potent and selective immunoproteasome inhibitor with minimal cross-reactivity for the constitutive proteasome in a cell-free assav Size 5 mg 25 mg

S7049 Oprozomib (ONX 0912)

Oprozomib (ONX 0912) is an orally bioavailable inhibitor for CT-L activity of 20S proteasome β5/LMP7 with IC₅₀ of 36 nM/82 nM. Phase 1/2

Size 5 mg 50 mg 10 mM/1 mL

S3017 Aspirin

Aspirin is a salicylate, and irreversible COX1 and COX2 inhibitor, used as an analgesic to relieve minor aches and pains, as an antipyretic to reduce fever, and as an anti-inflammatory medication.

Size 50 mg 1 g 5 g 10 mM/1 mL

the	Innibi	tory Selectivity					
L A	Inhibitor Name	DUB	USP/UBP	UCH	Other		
0	PR-619		++ EC50: 8.23 μM	+++ EC50: 2.95 μM	JOSD2, SENP6 cor DEN1		
	P5091		++ IC50: 4.3 μM		1		
112	TCID			+++ IC50: 0.6 μM			
	LDN-57444			++++ IC50: 0.88 μM			
	IU1		+ IC50: 4.7 μM				
C .	P22077		+ EC50: 8.6 μM				
<i>t</i>	VLX1570	+ IC ₅₀ : ~10 μM					
ibits	ML323	++++ IC ₅₀ : 76 nM					
little	b-AP15			+++ IC ₅₀ : 2.1 μM			
0он	Degrasyn	1			Bcr-Abl		

Notes:

DUB Inhibitors

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

S7130 PR-619

PR-619) is a non-selective, reversible inhibitor of the deubiquiti	nylating
enzyme	es (DOBS) with EC50 of 1-20 µm in a cell-free assay.	
Size	25 mg	$\sum_{h \in \mathbb{N}} \sum_{n \in \mathbb{N}} \sum_{$

S7132 P5091 (P005091)

P5091 (P005091) is a selective and potent inhibitor of ubiquitin-specific protease 7 (USP7) with EC50 of 4.2 µM and the closely related USP47. Size 10 mg 50 mg

S7134 IU1

d'i

IU1 is a cell-permeable, reversible and selective proteasome inhibitor of human USP14 with IC50 of 4.7 µ M. 25-fold selective to IsoT. Size 10 mg 50 mg 0790

S7529 ML323

ML323 displays reversible, nanomolar inhibitory activity and excellent selectivity toward USP1/UAF1 with IC50 of 76 nM. Size 5 mg 25 mg

Que and

USP/UBP selective

S2243 Degrasyn (WP1130)

Degrasyn (WP1130) is a selective deubiquitinase (DUB: USP5, UCH-L1, USP9x, USP14, and UCH37) inhibitor and also suppresses Bcr/Abl, also a JAK2 transducer (without affecting 20S proteasome) and activator of transcription (STAT).

----- Page 51

S8288 VLX1570

VLX1570 is a competitive inhibitor of proteasome DUB activity, with an IC₅₀ of ~10 µM in vitro. Size 5 mg

p97 / E2 Conjugating / E1 Activating / E3 Ligase

E3 Ligase Inhibitors Activator | Antagonists

E3 Ligase Inhibitors

S1193 Thalidomide

100-00

Thalidomide was introduced as a sedative drug, immunomodulatory agent and also is investigated for treating symptoms of many cancers. Thalidomide inhibits an E3 ubiquitin ligase, which is a CRBN-DDB1-Cul4A complex. Page 59

S2781 RITA (NSC 652287)

RITA (NSC 652287) induces both DNA-protein and DNA-DNA cross-links with no detectable DNA single-strand breaks, and also inhibits MDM2-p53 interaction by targeting p53.

S7892 Avadomide (CC-122)

Avadomide(CC-122), a new chemical entity termed pleiotropic pathway modifier, is a novel agent for Diffuse large B-cell lymphoma(DLBCL) with antitumor and immunomodulatory activity. Its molecular target is the protein cereblon (CRBN), a substrate receptor of the cullin ring E3 ubiquitin ligase complex CRL4CRBN.

Size 2 mg 5 mg 25 mg



X

E1 Activating Inhibitor

E2 Conjugating Inhibitor

S7129 PYR-41

p97 Inhibitor

S7285 NMS-873

(IC₅₀s >10 µM).

Size 5 mg 50 mg

S2913 BAY 11-7082

components of the ubiquitin system.

PYR-41 is the first cell-permeable inhibitor of ubiquitin-activating enzyme E1, with no activity at E2. Size 10 mg 25 mg 100 mg

NMS-873 is an allosteric and specific p97 inhibitor with IC₅₀ of 30 nM

that demonstrates potent selectivity for VCP/p97 compared to a panel of other AAA ATPases, Hsp90, and 53 additional analyzed kinases

BAY 11-7082 is a NF-κB inhibitor, inhibits TNFα-induced IκBα

phosphorylation with IC $_{50}$ of 10 μM in tumor cells. Also inhibiting

----- Page 105

E3 Ligase Activator

S2341 (-)-Parthenolide

(-)-Parthenolide, an inhibitor of the Nuclear Factor-KB Pathway, specifically depletes HDAC1 protein without affecting other class I/II HDACs; Also promotes the ubiquitination of MDM2 and activates p53 cellular functions.

Size 100 mg 250 mg

E3 Ligase Antagonists

S1061 Nutlin-3

Nutlin-3 is a potent and selective Mdm2 (RING finger-dependent ubiquitin protein ligase for itself and p53) antagonist with IC50 of 90 nM in a cell-free assay; stabilizes p73 in p53-deficient cells. ----- Page 59

S1172 JNJ-26854165 (Serdemetan)

JNJ-26854165 (Serdemetan) acts as a HDM2 ubiquitin ligase antagonist and also induces early apoptosis in p53 wild-type cells, inhibits cellular proliferation followed by delayed apoptosis in the absence of functional p53. Phase 1.

----- Page 58

96

Neuronal Signaling



S1575 RO4929097

S1528 LY2811376

nM and 5 nM, respectively. Phase 2.

S1262 Avagacestat (BMS-708163)

RO4929097 is a γ secretase inhibitor with IC $_{50}$ of 4 nM in a cell-free assay, inhibiting cellular processing of A β 40 and Notch with EC50 of 14

----- Page 93

Avagacestat (BMS-708163) is a potent, selective, orally bioavailable

 $\gamma\text{-secretase}$ inhibitor of Aβ40 and Aβ42 with IC $_{50}$ of 0.3 nM and 0.27 nM, demonstrating a 193-fold selectivity against Notch. Phase 2.

LY2811376 is the first orally available non-peptidic β -secretase (BACE1) inhibitor with ICso of 239 nM-249 nM, that act to decrease $A\beta$

secretion with EC50 of 300 nM, demonstrated to have 10-fold selectivity

towards BACE1 over BACE2, and more than 50-fold inhibition over

other aspartic proteases including cathepsin D, pepsin, or renin. Phase 1

----- Page 103

Gamma-secretase Inhibitors

Detailed product information is on page 92-93

Beta Amyloid Inhibitors

Inhibitory Selectivity

Inhibitor Name	Beta Amyloid	Other
DAPT (GSI-IX)	++ IC ₅₀ : 20 nM	
RO4929097	+++ ICso: 14 nM	γ secretase,γ secretase(ICN)
MK-0752	+++ ICso: 5 nM	
Avagacestat	++++ IC50: 0.3 nM	
LY2811376	+ EC50: ~300 nM	BACE1
EUK 134	1	

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

S2215 DAPT (GSI-IX)

DAPT (GSI-IX) is a novel y-secretase inhibitor, which inhibits Aß production with IC50 of 20 nM in HEK 293 cells. ----- Page 93

5-HT Receptor Inhibitor | Antagonist | Agonist | Modulator

5-HT Receptor Inhibitor

S1333 Fluoxetine HCI

Fluoxetine HCl is a selective serotonin-reuptake inhibitor (SSRI) at the neuronal membrane, used in the treatment of depression. to Que Size 25 mg 100 mg

S2459 Clozapine Clozapine is an atypical antipsychotic drug by acting as a 5-HT antagonist, used in the treatment of schizophrenia. Size 50 mg 10 mM/1 mL

5-HT Receptor Antagonist

5-HT1 selective

5-HT Receptor Agonist

S1436 Tianeptine sodium

Tianeptine sodium is a selective serotonin reuptake enhancer (SSRE), used for treating major depressive episodes. Size 10 mg 50 mg 100 mg 10 mM/1 mL



5-HT Receptor Modulator

S1283 Asenapine maleate

Asenapine maleate is a high-affinity antagonist of serotonin, norepinephrine, dopamine and histamine receptors, used for the treatment of schizophrenia and acute mania associated with bipolar disorder. 25 mg 100 mg Size

COX Inhibitors

Inhibitory Selectivity

Inhibitor Name	сох	COX-1	COX-2	Other
Celecoxib			++++ IC50: 40 nM	
Ibuprofen		+ ICso: 13 μΜ	+ IC ₅₀ : 370 μM	
Indomethacin		++ ICso: 0.28 μM	+ IC ₅₀ : 14 μM	
Rofecoxib			++++ IC50: 18 nM	
Diclofenac Sodium		+++ IC ₅₀ : 60 nM	+++ IC50: 200 nM	
Lumiracoxib		++ Κ.: 3.2 μM	+++ Ki: 60 nM	
Lornoxicam		++++ IC50: 5 nM	++++ IC50: 8 nM	
Naproxen Sodium		+ ICso: 8.7 μM	+ IC50: 5.2 μM	
Ketorolac		++ ICso: 1.23 μM	++ IC ₅₀ : 3.50 μΜ	
Valdecoxib			++++ IC50: 5 nM	
Tolfenamic Acid			+++ IC ₅₀ : 0.2 μM	
Amfenac Sodium Monohydrate		++ IC ₅₀ : 250 nM	+++ IC50: 150 nM	
Nimesulide			+ IC50: 26 μM	
Meclofenamate Sodium		++++ IC ₅₀ : 40 nM	+++ IC50: 50 nM	
Carprofen			++++ IC50: 30 nM	
Nepafenac		√		
Sulindac	1			
Meloxicam	1			
Aspirin		√		
Suprofen		√		
Piroxicam	√			
Ketoprofen		√		
Etodolac	√			
Ibuprofen Lysine	√			
Pranoprofen	√			
Asaraldehyde			√	
Zaltoprofen		1		
Acemetacin	1			
Bromfenac Sodium		√		
Nabumetone	1			
Niflumic acid			√	GABA receptor
Phenacetin	1			

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.cor 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

Neuronal Signaling

COX / GluR / Adrenergic Receptor

S1261	Celecoxib	Licer	nsed by Pfizer	COX-2 selective	S1638	lbuprofen	COX-1 selective
Celeco	xib is a selectiv	ve COX-2 inhibit	or with IC50 of 40 nM	I in Sf9 cells.	Ibuprofe and CO	n (Dolgesic) is an anti-infla X-2 with IC₅₀ of 13 uM and 3	mmatory inhibitor targeting COX-1 70 µM, respectively.
Vehi	icle Celecoxi	b 🔯 Nimesulide			Size	50 mg10 mM/1 mL	Light of
₹ ¹²	°°] –	Aorta	Product Citations (6):		S3043	Rofecoxib	COX-2 selective
Relative EL A expression (A 9 & 01			Blood, 2011, 118(22): 5 Br J Pharmacol, 2014, 	891-900 171(2): 498-508	Rofecox Size	tib is a COX-2 inhibitor with	C50 of 18 nM.
	0 Wild type	ЕС-А́МРК	Data from [Br J Pharm 171(2): 498-508] Celecoxib purchased fi	acol, 2014, rom Selleck			

and the second s

GluR Antagonist

(-)-MK 801 Maleate is a potent, selective and non-competitive NMDA

ADX-47273 is a potent and specific mGlu5 positive allosteric modulator (PAM) with EC50 of 0.17 µM, showing no activity at other mGlu

receptor antagonist with Kd of 37.2 nM in rat brain membranes.

S2876 (-)-MK 801 Maleate

Size 10 mg 50 mg 10 mM/1 mL

GluR Modulator

Size 5 mg 10 mg 10 mM/1 mL

S2690 ADX-47273

subtypes.

GluR Inhibitor | Agonist | Antagonist | Modulator

GluR Inhibitor

S2251 (-)-Huperzine A (HupA)

(-)-Huperzine A is a potent, highly specific and reversible inhibitor of acetylcholinesterase (AChE) with K of 7 nM, exhibiting 200-fold more selectivity for G4 AChE over G1 AChE. Also acts as an NMDA receptor antagonist. Phase 4.

Size 2 mg 5 mg 10 mg

GluR Agonist Signaling

S6001 LY404039

LY404039 is a potent agonist of recombinant human mGlu2/mGlu3 receptors with K of 149 nM/92 nM, shows >100-fold selectivity over ionotropic glutamate receptors, glutamate transporters, and other receptors. Phase 3.



Product Citations (3): Neuropharmacology, 2012, 62(7) PLoS One. 2011. 6(7): e22235

Data from [PLoS One, 2011, 6(7): LY404039 purchased from Selleck

Product Citations (2) J Clin Invest, 2013, 123(12): 5119-34 Antiviral Res. 2015, 120: 140-6

5119-341

from Selleck

Data from [J Clin Invest, 2013, 123(12);

Doxazosin Mesvlate (DOX) purchased

Adrenergic Receptor Inhibitor Agonist Antagonist

Adrenergic Receptor Inhibitor

S1324 Doxazosin Mesvlate

Doxazosin Mesylate, a quinazoline-derivative, selectively antagonizes postsynaptic α 1-adrenergic receptors, used in the treatment of high blood pressure and urinary retention associated with benign prostatic hyperplasia

Size 50 mg 10 mM/1 mL



Adrenergic Receptor Agonist S2566 Isoprenaline HCI

Isoprenaline HCI is a non-selective beta-adrenergic receptor agonist, used for the treatment of bradycardia and heart block. Size 50 mg 10 mM/1 mL

NMDA receptor selective

OD Ç

mGluR5 selectiv

Adrenergic Receptor Antagonist

S2038 Phentolamine Mesylate Phentolamine Mesylate is a nonselective alpha-adrenergic antagonist

with I		1 uM		cuve alpha-adre	nergie antagonia
with it	Joo 01 0.	i pivi.			
Size	50 mg	100 mg	10 mM/1 mL		(NH
					"n^ "

AChR Inhibitor | Agonist | Antagonist | Modulator

AChR Inhibitor

S2462 Donepezil HCI

Donepezil HCl is a specific and potent AChE inhibitor for bAChE and hAChE with IC50 of 8.12 nM and 11.6 nM, respectively.

Size 10 mg 50 mg 200 mg



AChR Agonist

S2455 Bethanechol chloride

Bethanechol chloride is a selective muscarinic receptor agonist without any effect on nicotinic receptors. Size 50 mg 10 mM/1 mL ...L.Nt a+

AChR Antagonist

S3005 Paroxetine HCI

Paroxetine HCl is an antidepressant drug of the SSRI type. Size 10 mg 50 mg 10 mM/1 mL

AChR Modulator

S2629 PNU-120596 (Nsc 216666)

PNU-120596 is a positive allosteric modulator of α7 nAChR with EC50 of 216 nM.

Size 10 mg 50 mg 200 mg 10 mM/1 mL



Product Citation (1): PLoS One, 2013, 8(8): e73581

> Data from [PLoS One, 2013, 8(8): o735811

PNU-120596 purchased from Sellect

Histamine Receptor Inhibitor Agonist | Antagonist

Histamine Receptor Inhibitor

S3208 Fexofenadine HCI (MDL 16455A) Fexofenadine HCI inhibits histamine H1 receptor with IC50 of 246 nM. Ĵ, Size 10 mg 50 mg 10 mM/1 mL Q.m

AChR / Histamine Receptor / Dopamine Receptor

Histamine Receptor Agonist

S1358 Loratadine

AChE selective

mAChR selective

<u>``</u> 200 M

nAChR selective

Ó

guu www.selleckchem.com

Loratadine is a histamine H1 receptor antagonist, used to treat allergies. Also acts as a selective inhibitor of B(0)AT2 with IC50 of 4 µM. Size 10 mg 50 mg 200 mg 10 mM/1 mL



Histamine Receptor Antagonist

S184	7 Clem	astine	Fumarate	H1 receptor selective
Clemantag	astine Fur onist with	narate (IC₅₀ of	Clemastine) is a selectiv 3 nM.	ve histamine H1 receptor
Size	50 mg	5 g	10 mM/1 mL	HO TOH CO

Dopamine Receptor Inhibitor | Agonist | Antagonists

Dopamine Receptor Inhibitor

S3163 Benztropine mesylate

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Benztropine mesylate is a dopamine transporter (DAT) inhibitor with
IC50 of 118 nM.
                                                        -B-0
Size 50 mg 10 mM/1 mL
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Dopamine Receptor Agonist

S2451 Amantadine HCI (1-adamantanamine HCI)

Amantadine HCI is used to treat or prevent infections of the respiratory tract caused by a certain virus.

Size 25 mg 100 mg 10 mM/1 mL

HCI

0H

Dopamine Receptor Antagonists

S1763 Quetiapine Fumarate

Quetiapine Fumarate is an atypical antipsychotic used in the treatment of schizophrenia, bipolar I mania, bipolar II depression, bipolar I depression and shows affinity for various neurotransmitter receptors including serotonin, dopamine, histamine, and adrenergic receptors.

Size 25 mg 50 mg 100 mg 10 mM/1 mL



S2456 Chlorpromazine HCI

Chlorpromazine HCl is a dopamine and potassium channel inhibitor with ICso of 6.1 and 16 μM for inward-rectifying K^{*} currents and time-independent outward currents. Size 50 mg 5 g 10 mM/1 mL



101



Opioid Receptor Agonist | Antagonist

Opioid Receptor Agonist

S2480 Loperamide HCI

Loperamide HCI is a selective µ-opioid receptor agonist opioid with K of 3.3 nM, 15-fold and 350-fold selective over the δ subtype and the μ versus the κ subtype of the opioid receptor, used against diarrhea resulting from gastroenteritis or inflammatory bowel disease. Size 50 mg 10 mM/1 mL



S2133 Gabapentin

S7071 (+)-Bicuculline

Size 50 mg 250 mg 10 mM/1 mL

S3066 Naloxone HCI

GABA Receptor Inhibitor | Activator | Agonist | Antagonist

GABA Receptor Inhibitor

S1168 Valproic acid sodium salt (Sodium valproate)

Valproic acid sodium salt (Sodium valproate) is a HDAC inhibitor by selectively inducing proteasomal degradation of HDAC2, used in the treatment of epilepsy, bipolar disorder and prevention of migraine headaches. ------ Page 22

neuropathic pain. Size 25 mg 100 mg

GABA Receptor Antagonist

of 2 µM, also blocks Ca(2+)-activated potassium channels.

GABA Receptor Agonist

Licensed by Pfizer

Gabapentin is a GABA analogue, used to treat seizures and

Opioid Receptor Antagonist

GABA Receptor Activator

S1969 Nefiracetam

Nefiracetam is a GABAergic, cholinergic, and monoaminergic neuronal systems enhancer for Ro 5-4864-induced convulsions. Phase 2.

Neuronal Signaling Size 50 mg 250 mg 10 mM/1 mL

P-qp Inhibitors | Modulator

Inhibitory Selectivity

Inhibitor Name	P-gp	Other
Zosuquidar 3HCI	++ K _i : 60 nM	
Tariquidar	+++ Kd: 5.1 nM	
Elacridar (GF120918)	1	BCRP

Notes:

1. For more details, such as half maximal inhibitory concentrations (IC50s) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

P-gp Inhibitors

S8028 Tariquidar

Tariquidar is a potent and selective noncompetitive inhibitor of P-alvcoprotein with Kd of 5.1 nM in CHrB30 cell line, reverses drug resistance in MDR cell Lines. Phase 3. and wat

Size 10 mg 50 mg 10 mM/1 mL



Selleck

LY2811376 is the first orally available non-peptidic β-secretase (BACE1) inhibitor with IC50 of 239 nM-249 nM, that act to decrease A β

secretion with EC50 of 300 nM, demonstrated to have 10-fold selectivity

towards BACE1 over BACE2, and more than 50-fold inhibition over

other aspartic proteases including cathepsin D, pepsin, or renin. Phase 1

P2 Receptor Inhibitor Antagonist

P2 Receptor Inhibitor

S1415 Clopidogrel

Clopid	logrel is	an oral,	, thienopyridine class antiplatelet agent.
Size	50 mg	200 mg	j.

P2 Receptor Antagonist

S4079 Ticagrelor

S7279 Suvorexant (MK-4305)

respectively. Phase 3.

Size 5 mg 50 mg

Ticagrelor is the first reversibly binding oral P2Y12 receptor antagonist with K of 2 nM.

Suvorexant (MK-4305) is a potent dual OX receptor antagonist with Ki

of 0.55 nM and 0.35 nM for OX1 receptor and OX2 receptor,

Size 50 mg 10 mM/1 mL



BACE Inhibitors

S1528 LY2811376

Ta2576+B-inhibitor

5 mg 10 mg 50 mg 10 mM/1 mL Tg2576+y-inhibito Product Citations (4):



Data from [J Biol Chem, 2014, 289(30): LY2811376 (β-inhibitor) purchased from Sellect

S8173 Verubecestat (MK-8931) Trifluoroacetat new

Verubecestat (MK-8931) is a potent and selective beta-secretase inhibitor and BACE1 protein inhibitor or Beta-site APP-cleaving enzyme 1 inhibitor

Size 5 mg 25 mg 100 mg



Neurona Signal

MT Receptor Agonist

OX Receptor Antagonist

S1259 Ramelteon

Ramelteon is a novel melatonin receptor agonist for human MT1 and MT2 receptors and chick forebrain melatonin receptors with K of 14 pM. 112 pM and 23.1 pM, respectively.

	т	argeted Biom	Table 9 arker Assay Resu	lts: Application	Set	
		Teratogenicity potential (µM)		Rodent in vivo test results*		
Compound	Cmax (µM)	o/c Ratio	Cell viability	Teratogenic ^b	Embryotoxic ^c	Cmax reference
6-Aminonicotinamide	NA	< 0.04	24.5	+4	۵.	NA
Abacavir	14.9	95.1	94.1	+	+	GlaxoSmithKline (2012)
Adefovir dipivoxil ^e	0.03	0.0015	0.02			Gilead Sciences (2012)
Amprenavir	15.1	236.9	259.5	+	+	GlaxoSmithKline (2005)
Artesunate	73.9	0.64	0.58	+1	+1	Miller et al. (2012)
Cidofovir ⁸	41.2	0.3	1.9			Gilead Sciences (2000)
Entacapone	3.9	6.7	127	+	-	Novartis Pharmaceuticals (2010)
Fluoxetine	0.04	25.1	23		+	Warner Chilcott (2013
Ramelteon ^h	0.02	34	>300	-		Karim et al. (2006)
Rosiglitazone	1.7	18.9	21.8		+	GlaxoSmithKline (2011)

Product Citation (1):

Ramelteon purchased from Selleck

KN-93 Phosphate ++ Ki: 0.37 µM Notes: 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

S7422 KN-62

KN-62 is a potent and specific inhibitor of Ca2+/calmodulin-dependent protein kinase II (CaMKII) with K_i of 0.9 µM; also a non-competitive antagonist of the purinergic receptor P2RX7 (IC50 = 15 nM). It is selective for CaMKII relative to PKA, PKC and MLCK, but inhibits CaMKI and CaMKIV equally well, The K_i value of KN-62 for CaMK V is 0.9......

oto pitti	0
Size 5 mg 25 mg	X
	Ån A
	25 - 3 - 8

S7423 KN-93 Phosphate

KN-93 Phosphate is a potent and specific inhibitor of Ca2+/calmodulindependent protein kinase II (CaMKII) with K of 0.37 $\mu\text{M},$ no remarkable inhibitory effects on APK, PKC, MLCK or Ca2+-PDE activities



S8366 CRT0066101

CRT0066101 is a small molecule PKD family specific inhibitor which specifically blocks PKD1/2 activity and does not suppress PKCq/ PKCβ/PKCε activity in multiple.

----- Page 60

1. For more details, such as half maximal inhibitory concentrations (IC50s) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com

NH125 KN-62 Ki: 0.9 µM

CaMK Inhibitors

Inhibitory Selectivity

Inhibitor Name CaMKI Othe CaMK L P2RX7 CaMK IV

Size 2 mg 10 mg 50 mg 10 mM/1 mL Ry

Birth Defects Res B Dev Reprod Toxicol, 2013, 98(4); 343-63

Data from [Birth Defects Res B Dev Reprod Toxicol, 2013, 98(4): 343-63]

P-gp Modulator

S1481 Zosuquidar (LY335979) 3HCI

S7772 Elacridar (GE120918)

Size 10 mg 50 mg 200 mg

Zosuguidar (LY335979) 3HCl is a potent modulator of P-glycoproteinmediated multi-drug resistance with K of 60 nM in a cell-free assay. Phase 3

Size 5 mg 10 mg 50 mg 10 mM/1 mL Product Citations (7)

J Lipid Res. 2014 56(1): 60-9 Aquatic Toxicology, 2014, 156C: 135-47

> Data from [Aquatic Toxicology, 2014, Zosuguidar 3HCI (ZSQ) purchased from



R

N



(+)-Bicuculline is a competitive antagonist of GABAA receptors with IC50





NF-kB Pathway



HDAC Inhibitors

Detailed product information is on page 19-23

NF-KB Inhibitors

Inhibitory Selectivity

Inhibitor Name	NF-ĸB	Other
QNZ (EVP4593)	++++ IC ₅₀ : 11 nM	TNF-α
JSH-23	++ IC ₅₀ : 7.1 μM	
SC75741	+++ EC50: 200 nM	
Sodium 4-Aminosalicylate	1	
Caffeic Acid Phenethyl Ester	1	
Sodium salicylate	√	
Andrographolide	1	

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

S4902 QNZ (EVP4593)

104

QNZ (EVP4593) shows potent inhibitory activity toward both NF-kB activation and TNF-α production with IC50 of 11 nM and 7 nM in Jurkat T cells, respectively. 9 mar

Size 5 mg 25 mg

S7351 JSH-23

Size

JSH-23 is an inhibitor of NF-KB transcriptional activity with IC50 of 7.1 uM in RAW 264.7 cell line. Jul O 5 mg 25 mg

S7414 Caffeic Acid Phenethyl Ester

Caffeic acid phenethyl ester is a potent and specific inhibitor of NF-kB activation, and also displays antioxidant, immunomodulatory and antiinflammatory activities. 20,000 Size 50 mg 200 mg

S7273 SC75741

Size 10 mg 50 mg



S3604 Triptolide (PG490)



Inhibitory	S	electiv	vity	/	
Inhibitor Name		lκB		ікк	Other
BAY 11-7082	++	IC ₅₀ : 10 µM			E2-conjugating enzymes
IKK-16			+++	IC50: 40 nM	
TPCA-1			++++	IC50: 17.9 nM	
BMS-345541			++	IC50: 0.3 µM	
SC-514			++	IC50: 3~12 µM	CDK2/CyclinA,AUR2,PRAK
Bay 11-7085	++	IC ₅₀ : 10 µM			
Rosmarinic acid			+	IC50: 12 µM	
MRT67307 HCI			+++	IC50: 160~190 nM	
PS-1145			+++	IC ₅₀ : 88 nM	
LY2409881			++++	IC50: 30 nM	
IMD 0354			1		
Bardoxolone Methyl			1		NF-ĸB,Nrf2
Mesalamine			1		
AZD3264			1		
WS6			1		EBP1
WS3			1		EBP1

Ir B/IKK Inhibitors

Notes:

1. For more details, such as half maximal inhibitory concentrations (IC50s) and working concentrations of each inhibitor, please visit the website of www.selleckchem.co 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

S2913 BAY 11-7082

BAY 11-7082 is a NF-κB inhibitor, inhibits TNFα-induced IκBα phosphorylation with IC50 of 10 µM in tumor cells. Also inhibiting components of the ubiquitin system

Size 10 mg 50 mg 10 mM/1 mL



S2824 TPCA-1

TPCA-1 is an inhibitor of IKK-2 with IC50 of 17.9 nM in a cell-free assay, inhibits NF-KB pathway, exhibits 22-fold selectivity over IKK-1.



S2882 IKK-16 (IKK Inhibitor VII)

IKK-16 (IKK Inhibitor VII) is a selective IkB kinase (IKK) inhibitor for IKK-2, IKK complex and IKK-1 with IC50 of 40 nM, 70 nM and 200 nM, respectively

Size 10 mg 50 mg 10 mM/1 mL

S2864 IMD 0354

IKB selective

-0+

IKK selective



IMD 0354 is an IKK β inhibitor and blocks IkB α phosphorylation in NF-k B pathway ψ

Size 5 mg 10 mg 50 mg 10 mM/1 mL

IKK selective

S8078 Bardoxolone Methyl

Bardoxolone Methyl is an IKK inhibitor, showing potent proapoptotic and anti-inflammatory activities; Also a potent Nrf2 activator and nuclear factor-κB (NF-κB) inhibitor.



S7352 Bay 11-7085

BAY 11-7085 is an irreversible inhibitor of TNF α -induced I κ B α phosphorylation with IC50 of 10 µM. ,0%~ Size 10 mg 25 mg

S8044 BMS-345541

BMS-345541 is a highly selective inhibitor of the catalytic subunits of IKK-2 and IKK-1 with IC50 of 0.3 µM and 4 µM in cell-free assays, respectively.

Size 5 mg 25 mg

S1274 BX-795

BX-795 is a potent and specific PDK1 inhibitor with IC50 of 6 nM, 140and 1600-fold more selective for PDK1 than PKA and PKC in cell-free assays, respectively. Meanwhile, in comparison to GSK3ß more than 100-fold selectivity observed for PDK1.



NOD1 Inhibitor

S2863 ML130 (Nodinitib-1)

ML130 (Nodinitib-1) is a potent and selective inhibitor of NOD1 with IC50 of 0.56 µM, inhibits NF-kB activation, exhibits 36-fold selectivity over NOD2.

Size 5 mg 25 mg 50 mg 10 mM/1 mL



IkB/IKK / NOD1

IrR selective

IKK selective

CLⁱL_n_m

IKK selective

NF-kB



Cannabinoid Receptor Agonist | Antagonist

Cannabinoid Receptor Agonist

S1544 AM1241

AM1241 is a selective cannabinoid CB2 receptor agonist with Ki of 3.4 nM, exhibits 82-fold selectivity over CB1 receptor. Size 2 mg 10 mg 25 mg 10 mM/1 mL ∞_{L}



Cannabinoid Receptor Antagonist

S3021 Rimonabant

Rimonabant is a selective antagonist of CB1 with IC50 of 13.6 nM and EC50 of 17.3 nM in hCB1 transfected HEK 293 membrane. Size 10 mg 50 mg 100 mg 10 mM/1 mL

Endothelin Receptor Antagonist

S4220 Bosentan

Bosentan is an endothelin (ET) receptor antagonist for ET-A and ET-B with K of 4.7 nM and 95 nM, respectively. Size 50 mg

S1P Receptor Inhibitor Antagonist | Modulator

S1P Receptor Inhibitor

S7177 PF-543

PF-543, a novel sphingosine-competitive inhibitor of SphK1, inhibits SphK1 with IC50 and Ki of 2.0 nM and 3.6 nM, exhibits >100-fold selectivity over the SphK2 isoform. andras

Size 10 mM/1 ml

S1P Receptor Antagonist

S5002 Fingolimod (FTY720) HCI



S1P Receptor Modulator

S7179 BAF312 (Siponimod)

BAF312 (Siponimod) is a next-generation S1P receptor modulator, selective for S1P1 and S1P5 receptors with EC50 of 0.39 nM and 0.98 nM, exhibits >1000-fold selectivity over S1P2, S1P3 and S1P4 receptors. Phase 3.

Size 5 mg 25 mg 100 mg

CR1 selective



GPCR

and

G

Protein

SGLT Inhibitors

S1548 Dapagliflozin

Dapagliflozin is a potent and selective hSGLT2 inhibitor with EC50 of 1.1 nM, exhibiting 1200-fold selectivity over hSGLT1. Phase 4. 5 mg 10 mg 50 mg 10 mM/1 mL Size



SGLT2 selective

S2760 Canagliflozin Canagliflozin is a highly potent and selective SGLT2 inhibitor for hSGLT2 with IC50 of 2.2 nM in a cell-free assay, exhibits 413-fold selectivity over hSGLT1. mar o

Size 5 mg 10 mg 10 mM/1 mL

SGLT2 selective

S8022 Empagliflozin (BI 10773) Empagliflozin (BI-10773) is a potent and selective SGLT-2 inhibitor with IC50 of 3.1 nM, exhibiting >300-fold selectivity over SGLT-1, 4, 5 and 6. Phase 3.

Size 5 mg 25 mg 10 mM/1 mL

LPA Receptor Antagonist

S1315 Ki16425

Ki16425 is a competitive, potent and reversible antagonist to LPA1, LPA2 and LPA3 with K of 0.34 µM, 6.5 µM and 0.93 µM, in RH7777 cell lines, respectively, shows no activity at LPA4, LPA5, LPA6. Size





Data from [J Bone Miner Metab, 2014, 10 1007/s00774-014-0607-51 Ki16425 purchased from Sellect

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CaSR Antagonist

Size 10 mg 100 mg 10 mM/1 mL

S2633 NPS-2143

of hyperparathyroidism.

Protein

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GPCR and

NPS-2143 is a novel potent and selective antagonist of Ca(2+) receptor with IC50 of 43 nM. Size 10 mg 50 mg - Hart Kron

Vasopressin Receptor Antagonist

S2593 Tolvaptan

Tolvaptan is an orally effective nonpeptide arginine vasopressin V2 receptor antagonist with IC50 of 3 nM, used to treat hyponatremia

Size 10 mg 50 mg 10 mM/1 mL

CXCR Antagonists

S8030 Plerixafor (AMD3100)

Plerixafor (AMD3100) is a chemokine receptor antagonist for CXCR4 and CXCL12-mediated chemotaxis with ICso of 44 nM and 5.7 nM in cell-free assays, respectively

Size 5 mg 10 mg 50 mg

S7651 SB225002

108

SB225002 is a potent, and selective CXCR2 antagonist with IC50 of 22 nM for inhibiting interleukin IL-8 binding to CXCR2, > 150-fold selectivity over the other 7-TMRs tested. an Orly

Size 10 mg 50 mg 200 mg

	Ag	onist Anta	agonist	
)	Ade	enosine Re	ceptor Inhil	oitor
	S8314	5-lodotubercidin	new	
	5-lodotu	ibercidin is a potent ad	enosine kinase inhibitor v	with IC50

10 mg 50 mg 100 mg 10 mM/1 mL



Adenosine Receptor Agonist

S2153 CGS 21680 HCI

CGS 21680 HCI is an adenosine A2 receptor agonist with IC50 of 22 nM. exhibits 140-fold over A1 receptor. Size 5 mg 25 mg 50 mg 10 mM/1 mL

Adenosine Receptor Antagonist

S2790 Istradefylline Istradefylline is a selective adenosine A2A receptor (A2AR) antagonist with Ki of 2.2 nM. Phase 3.

Size 5 mg 25 mg 10 mM/1 mL

Endocrinology and Hormones

Growth Factors RTKs GPCE IGE-1 IL-6R BEZ235 GDC-0941 LY294002 HS-173 (p110α) TGX-221 (p110β) CZC24832 (p110γ) CAL-101 (p110δ) Dahrafer Pan-Akt Inhibi MK-2206 GDC-0879 (B-Raf) GW5074 (C-Raf) Perifosine Endocrinology & A-674563 (Akt1) CCT128930 (Akt2) Inha Roducta Dutasteride Enzalutam Bicalutami Hormones Galeteron AZD3514 Vasodilation Raloxifene Tamoxifen ntiapoptosis Andarine Hexestrol Erteberel Eet. Gene Expression

Opioid Receptor Agonist | Antagonist

Detailed product information is on page 102

5-alpha Reductase Inhibitor | Antagonist

5-alpha Reductase Inhibitor

5-alpha Reductase Antagonist

S1197 Finasteride Finasteride is a potent, reversible inhibitor of the rat type 1 5 alphareductase with Ki of 10.2 nM, used in the treatment of benign prostatic hyperplasia (BPH) and male pattern baldness (MPB). Size 100 mg 200 mg

S1972 Tamoxifen Citrate Tamoxifen Citrate is an antagonist of the estrogen receptor by competitive inhibition of estrogen binding.

Estrogen/progestogen Receptor Inhibitor | Agonists | Antagonists | Chemical | Modulators

Ô. www.selleckchem.com

Estrogen/progestogen Receptor Estrogen/progestogen Receptor Inhibitor

S4285 Ospemifene

Ospemifene is a non-hormonal selective estrogen receptor modulator (SERM), used for the treatment of dyspareunia. \cap Size 25 mg 100 mg

Agonists

S2567 Medroxyprogesterone acetate

Medroxyprogesterone acetate is a progestin, a synthetic variant of the human hormone progesterone and a potent progesterone receptor agonist. Size 50 mg 10 mM/1 mL



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Adenosine Receptor Inhibitor

Estrogen/progestogen Receptor / Androgen Receptor

S2314 Kaempferol

Kaempferol, a natural flavonol, functions as an $\mathsf{ERR}\alpha$ and $\mathsf{ERR}\gamma$ inverse agonist. It inhibits topoisomerase I catalyzed DNA religation and may also inhibit the activity of fatty acid synthase.

Size 50 mg 200 mg

Tamoxifen Citrate is an antagonist of the estrogen receptor by competitive inhibition of estrogen binding. Size 250 mg 10 mM/1 mL

Chemical

S1251 Dienogest

Modulators

Size 10 mg 50 mg

S1972 Tamoxifen Citrate



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Estrogen/progestogen Receptor Antagonists

S1191 Fulvestrant

Fulvestrant is an estrogen receptor (ER) antagonist with IC50 of 0.94 nM in a cell-free assav.



S2606 Mifepristone

Mifepristone is a remarkably active antagonist of progesterone receptor and glucocorticoid receptor with IC50 of 0.2 nM and 2.6 nM, respectively.

Size 50 mg 200 mg 10 mM/1 mL





Androgen Receptor Inhibitor | Agonist | Antagonists | Modulator

Androgen Receptor Inhibitor

S2840 ARN-509

ARN-509 is a selective and competitive androgen receptor inhibitor with IC50 of 16 nM in a cell-free assay, useful for prostate cancer treatment. Phase 3.

______ 10 mg _____ 10 mM/1 mL Size 5 mg



Androgen Receptor Agonist

S2604 Dehydroepiandrosterone (DHEA)

Dehydroepiandrosterone is an important endogenous steroid hormone, which is an androgen receptor antagonist and an estrogen receptor agonist. Size 10 mg



Androgen Receptor Antagonists

S2803 Galeterone

Galeterone is a selective CYP17 inhibitor and androgen receptor (AR) antagonist with IC50 of 300 nM and 384 nM, respectively, and is a potent inhibitor of human prostate tumor growth. Phase 2. <u>8</u> dgⁱ Size 5 mg 25 mg 50 mg 10 mM/1 mL

S1190 Bicalutamide

Bicalutamide is an androgen receptor (AR) antagonist with IC50 of 0.16 μM. Size

200 mg 10 mM/1 mL 50 mg 100 mg



Int J Cancer, 2012, 131(6); E872-83 Data from [Oncogene, 2014, 10.1038/onc.2014.302] Bicalutamide purchased from Selleck

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Size

S1250 Enzalutamide (MDV3100)

Enzalutamide (MDV3100) is an androgen-receptor (AR) antagonist with IC50 of 36 nM in LNCaP cells.

Size 5 mg 10 mg 50 mg 10 mM/1 mL



Int J Cancer, 2012, 131(6): E872-83 Data from [Mol Cell Endocrinol, 2013. 365(1): 95-1071 WDV3100 purchased from Selleck

Androgen Receptor Modulator

S1174 MK-2866 (GTx-024)

MK-2866	(GTx-024)	is a	a selective	androgen	receptor	modulator
(SARM) w	ith Ki of 3.8	nM,	and is tiss	ue-selective	for anabo	olic organs.
Phase 3.						

Androgen Receptor / RAAS

RAAS Inhibitor | Antagonists

Inhibitory Selectivity

Inhibitor Name	AT1 receptor	AT2 receptor	ACE	Renin	RAAS
Aliskiren Hemifumarate				+++ IC50: 1.5 nM	
Candesartan	++++ IC50: 0.26 nM				
Losartan Potassium	+ IC ₅₀ : 20 nM				
Enalaprilat Dihydrate			+++ IC50: 1.94 nM		
Irbesartan	+++ IC50: 1.3 nM				
PD123319		+ IC ₅₀ : 34 nM			
Perindopril Erbumine			+++ IC50: 1.05 nM		
Candesartan Cilexetil					++++ IC50: 0.26 nM
Ramipril			++ IC50: 5 nM		
Captopril			+ IC50: 6 nM		
Azilsartan Medoxomil	++ IC50: 2.6 nM				
Imidapril HCI			++ IC50: 2.6 nM		
Eprosartan Mesylate	++++ Kd: 0.83 nM				
Azilsartan	++ IC50: 2.6 nM				
Telmisartan		1			
Valsartan		1			
Benazepril HCI			√		
Enalapril Maleate			√		
Olmesartan Medoxomil	√				
Cilazapril Monohydrate			√		
Lisinopril			√		
Moexipril HCI			1		

110



contraception and the treatment of endometriosis.

Size 10 mg 100 mg 1 g 10 mM/1 mL

Estrogen/progestogen Receptor

Dienogest is an orally active synthetic progesterone, used for

Estrogen/progestogen Receptor

selective estrogen receptor (ER) modulator that is widely used in the

therapeutic and chemopreventive treatment of breast cancer.

Inhibitory Selectivity

Inhibitor Name	AT1 receptor	AT2 receptor	ACE	Renin	RAAS
Temocapril			√		
Temocapril HCI			1		
Quinapril HCI			√		
LCZ696					1
Fosinopril Sodium			√		

Notes:

Endocrinology & Hormones

1. For more details, such as half maximal inhibitory concentrations (ICcos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

RAAS Inhibitor

S1235 Letrozole Letrozole is a third generation inhibitor of aromatase with IC_{50} of 0.07



Aliskiren Hemifumarate is a direct renin inhibitor with IC50 of 1.5 nM.



RAAS Antagonists

S1359 Losartan Potassium (DuP 753)

Losartan Potassium is an angiotensin II receptor antagonist, con with the binding of angiotensin II to AT1 receptors with IC50 of 20 Size 50 mg 10 mM/1 mL

S7678 LCZ696

LCZ696, consisting of valsartan and sacubitril in 1:1 molar ratio orally bioavailable, dual-acting angiotensin receptor-neprilysin in (ARNi) for hypertension and heart failure. Phase 3.

Size 5 mg 25 mg 100 mg

o of 1.5 nM.	hormone (LH), follicle-stimulating horm	none (FSH), or androstenedione
ÇiXin j	in clinical studies.	blyte excretion or thyroid function
Lezan .	Size 25 mg 50 mg 200 mg 10 mM/	<u>1 mL</u> CLO
AT1 receptor selective nist, competes 30 of 20 nM.	C C+T C+T+D	Carbon Control
jo Ray	S1196 Exemestane	Licensed by Pfizer
HO N	Exemestane is an aromatase inhibitor rat ovarian aromatase with IC50 of 30 n	, inhibiting human placental and M and 40 nM, respectively.
olar ratio, is an rilysin inhibitor	<u>Size 10 mg 50 mg 100 mg 10 mM/</u>	<u>1 mL</u>
NN SPEC		

-20 nM in cell-free assays. It has no effect on the plasma levels of 17

α-OH progesterone, thyroid-stimulating hormone (TSH), luteinizing

GPR Agonist | Antagonist

TAK-875 is a selective GPR40 agonist with EC50 of 14 nM in human

Aromatase Inhibitors

Inhibitory Selectivity

	*
Inhibitor Name	Aromatase
Letrozole	++++ IC50: 0.07-20 nM
Anastrozole	+++ IC ₅₀ : 15 nM
Exemestane	+++ ICso: 30 nM
Formestane	++ ICso: 80 nM
Aminoalutothimido	+ IC: 10 uM

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICros) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

S1188 Anastrozole

Anastrozole is a third-generation nonsteroidal selective aromatase inhibitor. It may offer greater selectivity compared with other aromatase inhibitors, being without any intrinsic endocrine effects and with no apparent effect on the synthesis of adrenal steroids. NON-

Size 10 mg 100 mg 1 g 10 mM/1 mL



GPR Antagonist

S7263 AZD1981

Excellent Validation, Technical Support and Prompt Delivery

GPR Agonist

S2637 TAK-875



Transmembrane Transporters



GABA Receptor Inhibitor | Activator | Agonist | Antagonist

Detailed product information is on page 102

P-qp Inhibitors | Modulator

Detailed product information is on page 102

Calcium Channel Inhibitor | Activator | Antagonist

Calcium Channel Inhibitor

S2403 Tetrandrine

Tetrandrine, a bis-benzylisoquinoline alkaloid derived from Stephania tetrandra, is a calcium channel blocker. Size 100 mg 10 mM/1 mL







S2482 Manidipine 2HCI



S2050 Strontium Ranelate

Strontium Ranelate is a strontium(II) salt of ranelic acid for (-)desmethoxyverapamil binding to calcium channel with IC50 of 0.5 mM. Size 50 mg 200 mg





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Calcium Channel Antagonist

Sodium Channel Inhibitor Antagonist

Inhibitory Selectivity

Inhibitor Name	Sodium Channel	Other
Oxcarbazepine	+ ICso: 160 μM	
Riluzole	1	NMDA receptor, Glutamate release
Amiloride HCI	1	T-type calcium channel,uPA
Rufinamide	1	
Zonisamide	1	
Phenytoin Sodium	1	
Amiloride HCI dihydrate	1	
Dronedarone HCI	1	Potassium channel, Calcium channel
Phenytoin	1	
Lamotrigine	1	5-HT (human platelets), 5-HT (rat brain synaptosomes)
Primidone	1	
Procainamide HCI	1	DNA methyltransferase
Digoxin	1	
Mexiletine HCI	1	
Benzocaine	1	
Tolperisone HCI	1	
Levobupivacaine HCI	1	
Dibucaine HCI	1	
Ibutilide Fumarate	1	
Vinpocetine	1	
Propafenone HCI	1	

Notes:

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1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com.

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

Sodium Channel Inhibitor

S4016 Quabain

Ouabain is a selective Na*/K*, -ATPase inhibitor, binds to $\alpha 2$ / $\alpha 3$ subunit with K of 41 nM/15 nM. Size 50 mg 10 mM/1 mL HO-C-O-CAN ON

Sodium Channel Antagonist

S1828 Proparacaine HCI

Proparacaine HCl is a voltage-gated sodium channels antagonist with ED50 of 3.4 mM ~ join Size 50 mg 10 mM/1 mL

ATPase Inhibitors | Activator

ATPase Inhibitors

S1478 Oligomycin A

Oligomycin A is an inhibitor of ATP synthase, inhibits oxidative phosphorylation and all the ATP-dependent processes occurring on the coupling membrane of mitochondria.

Size 5 mg 10 mg 25 mg 10 mM/1 mL

S7046 Brefeldin A

Brefeldin A is a lactone antibiotic and ATPase inhibitor for protein transport with IC50 of 0.2 µM in HCT 116 cells, induces cancer cell differentiation and apoptosis. Size 5 mg 25 mg 100 mg 10 mM/1 mL



S7099 (-)-Blebbistatin

Size 10 mg 25 mg 10 mM/1 mL

(-)-Blebbistatin is a cell-permeable inhibitor for non muscle myosin II ATPase with IC50 of ~2 µM, does not inhibit myosin light chain kinase, inhibits contraction of the cleavage furrow without disrupting mitosis or contractile ring assembly. τ, in

S8101 CB-5083

Size 5 mg 25 mg

CB-5083 is a potent, selective, and orally bioavailable p97 AAA ATPase inhibitor with IC50 of 11 nM. Phase 1.



ATPase Activator

S2623 Omecamtiv mecarbil (CK-1827452)

Omecamtiv mecarbil (CK-1827452) is a specific cardiac myosin activator and a clinical drug for left ventricular systolic heart failure. Phase 2.





Omecamtiv mecarbil (OM) purchased from Selleck

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Potassium Channel Inhibitor Activator | Antagonist

Potassium Channel Inhibitor

S2456 Chlorpromazine HCl

Chlorpromazine HCl is a dopamine and potassium channel inhibitor with IC50 of 6.1 and 16 µM for nward-rectifying K* currents and time-independent outward currents. ----- Page 101

Potassium Channel Activator

S1971 Nicorandil

Nicorandil is a potassium channel activator, and stimulates guanylate cyclase to increase formation of cyclic GMP (cGMP). Size 50 mg 10 mM/1 mL An y Potassium Channel Antagonist CFTR Modulators

S3151 Gliquidone

Gliquidone is an ATP-sensitive K* channel antagonist with IC50 of 27.2 nM Size 50 mg 10 mM/1 mL

Bafilomycin A1 is a vacuolar H*-ATPase inhibitor with IC50 of 0.44 nM.



VX-809 (Lumacaftor) acts to correct CFTR mutations common in cystic fibrosis by increasing mutant CFTR (F508del-CFTR) maturation, EC50 of 0.1 µM in fisher rat thyroid cells. Phase 3.





VX-809 purchased from Selleck

S7059 VX-661

Size

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VX-661 is a second F508del CFTR corrector and is believed to help CFTR protein reach the cell surface. Phase 2. 5 mg 50 mg 10 mM/1 mL Size



CFTR Inhibitors | Activator | **Modulators**

Proton Pump Inhibitor

CFTR Inhibitors

S1413 Bafilomycin A1 (Baf-A1)

Size 1 mg

S6003 Ataluren (PTC124)

Ataluren (PTC124) selectively induces ribosomal read-through of premature but not normal termination codons, with EC50 of 0.1 µM, may provide treatment for genetic disorders caused by nonsense mutations (e.g. CF caused by CFTR nonsense mutation). Phase 3.

Size 10 mg 50 mg 100 mg 10 mM/1 mL



S7139 CFTRinh-172

CFTRinh-172 is a voltage-independent, selective CFTR inhibitor with K of 300 nM, showing no effects on MDR1, ATP-sensitive K* channels, or a series of other transporters.

Size 10 mg 50 mg

CFTR Activator

S1144 Ivacaftor (VX-770)

Ivacaftor (VX-770) is a selective potentiator of CFTR targeting G551D-CFTR and F508del-CFTR with EC50 of 100 nM and 25 nM in fisher rat thyroid cells, respectively,

Size	5 mg	10 mg	50 mg	10 mM/1 mL



CRM1 Inhibitors

S7252 Selinexor (KPT-330) Selinexor (KPT-330) is an orally bioavailable selective CRM1 inhibitor Phase 2

Size 5 mg 50 mg

S7125 KPT-185 O. S. in

KPT-185 is a selective CRM1 inhibitor that induces growth inhibition and apoptosis in a panel of NHL cell lines with a median IC₅₀ ~25 nM. Size 10 mg 50 mg

S7551 Piperlongumine

Piperlongumine, a natural alkaloid from Piper longum L., increases the level of reactive oxygen species (ROS) and selectively kills cancer cells. It is a direct TrxR1 inhibitor with suppressive activity against gastric cancer and a novel inhibitor of CRM1: also an inhibitor of PI3K/Akt/mTOR in human breast cancer cells

Size	10 mg	50 mg	200 mg	ć

S8397 Eltanexor (KPT-8602)

Eltanexor(KPT-8602) is a second-generation, orally bioavailable XPO1 inhibitor with IC50 values of 20-211 nM in 10 AML lines after 3 days exposure.

Size 2 mg 5 mg 25 mg



TRPV Antagonist

S7115 AMG-517

AMG-517 is a potent and selective TRPV1 antagonist, and antagonizes capsaicin, proton, and heat activation of TRPV1 with IC₅₀ of 0.76 nM. 0.62 nM and 1.3 nM, respectively Size 5 mg 25 mg 10 mM/1 mL



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Data from [J Clin Invest, 2013, 124(1); 111-61 Ataluren purchased from Selleck



Product Citations (12): J Clin Invest, 2013, 124(1): 111-6 Hum Mol Genet, 2015, 24(4): 972-86

Metabolism



Metabolism

Detailed product information is on page 74-75

HSP (e.g. HSP90) Inhibitors

PPAR Inhibitor | Activator | Agonists | Antagonist

PPAR Inhibitor

S2871 T0070907

116

Modulator

T0070907 is a potent and selective PPARy inhibitor with IC $_{50}$ of 1 nM in a cell-free assay, with a >800-fold selectivity over PPARα and PPARδ. Size 5 mg 25 mg 50 mg 10 mM/1 mL States.

PPAR Activator

S8029 WY-14643 (Pirinixic Acid)

WY-14643 (Pirinixic Acid) is a potent peroxisome proliferator and activator of PPARa with EC50 of 1.5 µM. Size 50 mg 250 mg 10 mM/1 mL $Q, \dot{Q},$

PPAR Agonists

S2505 Rosiglitazone maleate

Rosiglitazone, a member of the thiazolidinedione class of antihyperglycaemic agents, is a high-affinity selective agonist of the peroxisome proliferator-activated receptor-y with IC50 of 42 nM. Size 25 mg 200 mg 1 g 10 mM/1 mL n de la Constance



Data from [Stroke, 2013, 44(12): 3498-5081 Rosiglitazone maleate (RSG) purchased from Selleck

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S2556 Rosiglitazone

Rosiglitazone is a potent antihyperglycemic agent and a potent thiazolidinedione insulin sensitizer with IC₅₀ of 12, 4 and 9 nM for rat, 3T3-L1 and human adipocytes, respectively. Rosiglitazone is a pure ligand of PPAR-gamma, and has no PPAR-alpha-binding action.



PPAR Antagonist

S2915 GW9662 (TIMTEC-BB, SBB006523

GW9662 is a selective PPAR antagonist for PPARy with IC₅₀ of 3.3 nM, with at least 100 to 1000-fold functional selectivity in cells with PPARy versus PPARα and PPARδ.

Size 10 mg 25 mg 50 mg 10 mM/1 mL

S2262 Apigenin

S2246 Abiraterone Acetate (CB7630)

Galeterone is a selective CYP17 inhibitor and androgen receptor (AR) antagonist with IC50 of 300 nM and 384 nM, respectively, and is a potent inhibitor of human prostate tumor growth. Phase 2.

P450 (e.g. CYP17) Inhibitors

S1123 Abiraterone (CB-7598) CYP17 selective Abiraterone is a potent CYP17 inhibitor with IC50 of 2 nM in a cell-free

Size 5 mg 25 mg

assay.



PDE Inhibitors

Inhibitory Soloctivity

Inhibitor Name	PDE	PDE1	PDE2	PDE3	PDE4	PDE5	PDE6	PDE10A	Other
Roflumilast					++++ IC ₅₀ : 0.7 nM				
Sildenafil Citrate						+++ IC50: 3.5 nM	+++ IC50: 33 nM		
Cilomilast					+++ IC50: 100 nM				
Tadalafil						+++ ICso: 1.8 nM			
Vardenafil HCI Trihydrate		+++ IC50: 180 nM				++++ ICso: 0.7 nM			
Pimobendan				++ IC ₅₀ : 0.32 μΜ					
GSK256066					++++ IC ₅₀ : 3.2 pM				
PF-2545920								++++ IC50: 0.37 nM	
Rolipram					++ ICso: 2.0 μM				
Cilostazol				++ IC50: 0.2 μM					
Milrinone			++ ICso: 5.2 μM	++ ICso: 2.1 μM					ATPase

PPAR / P450 (e.g. CYP17) / PDE



Abiraterone Acetate is an acetate salt form of Abiraterone which is a

PDE / Hydroxylase / Factor Xa

Inhibitory Selectivity

Inhibitor Name	PDE	PDE1	PDE2	PDE3	PDE4	PDE5	PDE6	PDE10A	Other
Avanafil						++++ IC50: 1 nM			
S- (+)-Rolipram					++ ICso: 0.75 μM				
Aminophylline	+ IC50: 0.12 mM								adenosine receptor
TAK-063								++++ IC50: 0.3 nM	
Deltarasin	+++ K _d : 38 nM								
Luteolin		+ Κι: 15.0 μΜ	+ Κι: 6.4 μΜ	+ Κι: 13.9 μΜ	+ Κι: 11.1 μΜ	+ Κι: 9.5 μΜ			
Icariin						++ IC50: 0.432 μM			
Anagrelide HCI	4								
Irsogladine	4								mAChR,AChR
Doxofylline	4								
Dipyridamole	4								
Dyphylline	4								

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsss) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com

PDE5 selective

PDF4 selectiv

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Notes

Excellent Validation, Technical Support and Prompt Delivery

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

S1431 Sildenafil Citrate

Sildenafil Citrate, a selective inhibitor of cyclic guanosine monophosphate (cGMP)-specific phosphodiesterase type 5 (PDE5), is a well-tolerated and highly effective treatment for erectile dysfunction.

Size 25 mg 50 mg 500 mg 10 mM/1 mL

ANK. veh Sil D+S D+S Dox D+S Product Citations (5): LC3/LC3/I Mol Pharmacol, 2014, 85(3); 408-19 p62 LAMP2 GAPDH J Cell Physiol, 2015, 10.1002/jcp.24961 ----Data from [Mol Pharmacol, 2014, 85(3): 6 h 24 h 408-191 Sildenafil Citrate (Sil) purchased from Selleck

S1512 Tadalafil (IC351)

Metabolism

Tadalafil is a PDE-5 inhibitor with IC50 of 1.8 nM in a cell-free assay. Tadalafil is at least 9000 times more selective for PDE5 than most of the other families of PDEs, with the exception of PDE11. It can partial inhibits PDE11.

Size 50 mg 100 mg 500 mg 10 mM/1 mL

S1430 Rolipram

The PDE4 selective inhibitor, rolipram, inhibited immunopurified PDE4B and PDE4D activities similarly, with IC $_{\rm 50}$ values of approx. 130 nM and 240 nM respectively; an anti-inflammatory agent.

Size 10 mg 25 mg 50 mg 10 mM/1 mL

S2320 Luteolin

Luteolin is a flavonoid found in Terminalia chebula which is a non-selective phisphodiesterase PDE inhibitor for PDE1-5 with Ki of 15.0 µM, 6.4 µM, 13.9 µM, 11.1 µM and 9.5 µM, respectively. Phase 2.

Size 10 mg 50 mg 200 mg 10 mM/1 mL

Hydroxylase Inhibitor Activator

Hydroxylase Inhibitor

S7483 DMOG (Dimethyloxalylglycine)

DMOG is an antagonist of α -ketoglutarate cofactor and inhibitor for HIF prolyl hydroxylase. Size 10 mM/1 mL j.

Hydroxylase Activator

Isotretinoin was developed to be used as a chemotherapy medication for the treatment of brain cancer, pancreatic cancer and more Size 50 mg 10 mM/1 mL

Factor Xa Inhibitors

Inhibitory Selectivity

Inhibitor Name	Factor Xa	Other
Rivaroxaban	++ IC50: 0.7 nM	Prothrombinase
Apixaban	++++ K _i : 0.08 nM	
Edoxaban	+++ Ki: 0.561 nM	

1. For more details, such as half maximal inhibitory concentrations (IC50s) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

S3002 Rivaroxaban (BAY 59-7939)

Rivaroxaban is a direct inhibitor of Factor Xa with K_i and IC_{50} of 0.4 nM and 0.7 nM in cell-free assays, respectively. It is selective for human factor Xa, for which it has >10 000-fold greater selectivity than for other biologically relevant serine proteases (IC₅₀ >20 µM). $(1-1)^{-1}(1-1$

Size 5 mg 10 mg 50 mg 10 mM/1 mL

S1593 Apixaban

Apixaban is a highly selective, reversible inhibitor of Factor Xa with Ki of 0.08 nM and 0.17 nM in human and rabbit, respectively. Size 10 mg 50 mg 10 mM/1 mL

DHFR Inhibitors

Inhibitory Selectivity

Inhibitor Name	DHFR	Other
Pemetrexed	++++ Ki: 7.2 nM	TS,GARFT
Methotrexate	+ ICso: 24 nM	
Pyrimethamine	++ IC ₅₀ : 15.4 nM	
Pemetrexed Disodium Hydrate	++++ Ki: 7.2 nM	TS,GARFT

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

S1135 Pemetrexed (LY-231514)

Pemetrexed is a novel antifolate and antimetabolite for TS, DHFR and GARFT with K of 1.3 nM, 7.2 nM and 65 nM, respectively.

Size 10 mg 50 mg 200 mg



Cancer Res, 2014, 74(21): 5948-54 Mol Cancer Res. 2013, 12(12): 2675-84 Data from [Mol Cancer Res, 2013,

Product Citations (4)

12(12): 2675-84] Pemetrexed purchased from Selleck

S1210 Methotrexate

Methotrexate (MTX), analog of folic acid, is a nonspecific inhibitor of the dihydrofolate reductase(DHFR) of bacteria and cancerous cells as well as normal cells. It forms an inactive ternary complex with DHFR and NADPH.

Size 100 mg 500 mg 10 g 10 mM/1 mL

Aminopeptidase Inhibitor

S1591 Bestatin

Bestatin is a potent aminopeptidase-B and leukotriene (LT) A4 hydrolase inhibitor, used in the treatment of acute myelocytic leukemia. Size 10 mg 25 mg 50 mg 100 mg

Dehydrogenase Inhibitors

Inhibitory Selectivity

nhibitor Name	Dehydrogenase
lycophenolate Mofetil	+++ ICso: 39 nM
GI-5198	++ ICso: 0.16 μM
IK-8245	++++ IC ₅₀ : 1 nM
nasidenib	++++ IC50: 12 nM
ICT-501	++ IC50: 40 nM
W033291	++++ Ki: 0.1 nM
/idofludimus	+ IC ₅₀ : 134 nM
GI-6780	+++ IC50: 23 nM
CPI-613	√
eflunomide	V
Disulfiram	V
rilostane	V
eriflunomide	√
PluriSIn #1 (NSC 14613)	V
mmonium Glycyrrhizinate	V
Simeracil	V
vosidenib (AG-120)	V
sovaleramide	V
Sossypol Acetate	<u>√</u>
noxolone	1
modin	1

Notes:

1. For more details, such as half maximal inhibitory concentrations (IC $_{\rm SO}s)$ and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

$= \sum_{m=1}^{n} \sum_{j=1}^{n} \sum_{i=1}^{n} \sum_{j=1}^{n} \sum_{j=1}^{n} \sum_{i=1}^{n} \sum_{j=1}^{n} \sum_{i=1}^{n} \sum_{j=1}^{n} \sum_{j=1}^{n} \sum_{i=1}^{n} \sum_$ S2776 CPI-613

CPI-613, a lipoate analog, inhibits mitochondrial enzymes pyruvate dehydrogenase (PDH) and α-ketoglutarate dehydrogenase in NCI-H460 cell line, disrupts tumor cell mitochondrial metabolism. Phase 2.

Size 5 mg 50 mg 10 mM/1 mL

S7185 AGI-5198 (IDH-C35)

AGI-5198 is the first highly potent and selective inhibitor of IDH1 R132H/R132C mutants with IC50 of 0.07 µM/0.16 µM.

Size 5 mg 25 mg

S2303 Gossypol Acetate

Gossypol Acetate is a polyphenolic aldehyde that permeates cells and acts as an inhibitor for several dehydrogenase enzymes such as lactate dehydrogenase, NAD-linked enzymes.

Size 100 mg 250 mg



S8206 Ivosidenib (AG-120)

Ivosidenib (AG-120) is an orally available inhibitor of isocitrate dehydrogenase type 1 (IDH1), with potential antineoplastic activity. Size 5 mg 25 mg



Metabo

60

Man Offer



Dehydrogenase / Procollagen C Proteinase / Carbonic Anhydrase / MAO / Phospholipase (e.g. PLA) / FAAH

S8205	Enasidenil) (AG-221)	new
Enaside inhibitor	nib (AG-221) of the IDH2	is a first-ir nutant en	n-class, oral, potent, reversible, sel zyme with IC50 of 12 nM.
Size	5 mg 25 mg	100 mg	í



Carbonic Anhydrase Inhibitors

Inhibitory Selectivity

Inhibitor Name	Carbonic Anhydrase	(Ai	Carbonic nhydrase I	Carbonic Anhydrase II	C: Anh	arbonic lydrase IV	C An	arbonic hydrase IX	Carbonic Anhydrase XII	Carbonic Anhydrase XII
Dorzolamide HCI		+	K _i : 6000 nM	++++ K _i : 1.9 nM	+++	K _i : 31 nM				
U-104							++	Ki: 45.1 nM	++++ Ki: 4.5 nM	
Tioxolone		++	Ki: 91 nM							
Brinzolamide				++++ IC50: 3.19 nM						
Methazolamide		++	K _i : 50 nM	+++ K _i : 14 nM	++	K _i : 36 nM				
Topiramate	٧									sodium channel, AMPA/kainate receptor, Calcium Channel
Dichlorphenamide	*									

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com.

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

S1438 Topiramate

Topiramate is a mutil-targeted inhibitor, including voltage-gated sodium channel and calcium channel, AMPA/kainate receptor and carbonic anhydrase, used to treat epilepsy.

Size 100 mg 10 mM/1 mL

MAO Inhibitor



Phospholipase (e.g. PLA)

agonist-induced Ca2+ increases in platelets and PMN.

U73122 is a potent phospholipase C (PLC) inhibitor, which reduces

Metabolism

Inhibitory Selectivity					
Inhibitor Name	MAO-A	MAO-B	MAO		
Safinamide Mesylate		++++IC50: 98 nM			
Rasagiline Mesylate	+++ IC50: 412 nM	++++IC50: 4.43 nM			
Moclobemide	+++ IC ₅₀ : 6.1 μΜ				
Sennoside A			++ IC ₅₀ : 17 μΜ		
Paeonol	+ IC50: 54.6 μM	++ IC50: 42.5 μM			

Notes:

Tranylcypromine HCI

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

S4246 Tranylcypromine (2-PCPA) HCI

Tranylcypromine (2-PCPA) HCl is a monoamine oxidase inhibitor, which inhibits CYP2A6 with K of 0.08 µM and 0.2 µM in cDNA-expressing microsomes and Human Liver Microsomes, respectively.

CAN H

FAAH Inhibitor

Size 5 mg 25 mg 100 mg

Inhibitor

S8011 U73122

S2631 URB597 (KDS-4103)

URB597 is a potent, orally bioavailable FAAH inhibitor with IC50 of 4.6 nM, with no activity on other cannabinoid-related targets. Phase 1. Size 5 mg 25 mg 100 mg 10 mM/1 mL $\mathrm{O}^{\mathrm{O}_{\mathrm{s}}\mathrm{i}_{\mathrm{p}}\mathrm{O}}$

OLO^{In}

IDO / Transferase / HMG-CoA Reductase

S1453 Tipifarnib (R115777)

Tipifarnib (R115777) is a potent and specific farnesyltransferase (FTase) inhibitor with IC50 of 0.6 nM, its anti-proliferative effects are most prominent in H-ras or N-ras mutant cells. Phase 3.



S2799 Daporinad (FK866, APO866)

Daporinad (FK866, APO866) effectively inhibits nicotinamide phosphoribosyltransferase (NMPRTase) with IC50 of 0.09 nM in a cell-free assay. Phase 1/2.

5 mg 10 mg 50 mg

S2821 RG108

 $\underset{N \in \mathbb{N}}{\overset{HO}{\longrightarrow}} \underset{\underline{n} \in \mathbb{N}}{\overset{N}{\longrightarrow}} \underset{\underline{n} \in \mathbb{N}}{\overset{N}}{\underset{\underline{n} \in \mathbb{N}}}{\overset{N}{\underset{\underline{n} \in \mathbb{N}}}{\overset{N}{\underset{\underline{n} \in \mathbb{N}}}$

July a

RG108 is an inhibitor of DNA methyltransferase with IC50 of 115 nM. does not cause trapping of covalent enzymes. Page 32

S7910 Epacadostat (INCB024360)

IDO-mediated immune suppression. Phase 2.

-dioxygenase) inhibitor with IC50 of 67 nM. Phase 2.

IDO Inhibitors

TDO

S7587 INCB024360 analogue

Size 5 mg 25 mg 100 mg

S7756 Indoximod (NLG-8189)

S7111 NLG919

Size 10 mg 50 mg

2.0×106

1.5-109

1.0×10

5.0×10

Epacadostat (INCB024360) is a potent and selective indoleamine 2,3-dioxygenase (IDO1) inhibitor with IC₅₀ of 10 nM and displays high selectivity over other related enzymes such as IDO2 or tryptophan 2,3-dioxygenase (TDO).

NLG919 is a potent IDO (indoleamine-(2,3)-dioxygenase) pathway

Product Citation (1):

J Biomol Screen, 2014, 19(9); 1266-74

Data from [J Biomol Screen, 2014, 19(9): 1266-74] NLG919 purchased from Selleck

inhibitor with K/EC50 of 7 nM/75 nM in cell-free assays. Phase 1.

- S7111

- NTRC 0672-0

INCB024360 analogue is a potent, competitive IDO1 (indoleamine-(2,3)

Indoximod (NLG-8189), a methylated tryptophan, acts as an IDO

(indoleamine-(2,3)-dioxygenase) pathway inhibitor, and reverses

Size 5 mg 25 mg

Size 50 mg 200 mg

Transferase Inhibitors

Inhibitory Selectivity

Tipifamib +++ IC ₅₀ : 0.6 nM Lonafamib +++ IC ₅₀ : 1.9-5.2 nM Daporinad (FK866, APO866) ++++ K: 0.4 nM A922500 +++ IC ₅₀ : 7-24 nM	
Lonafamilb +++ IC ₅₀ : 1.9~5.2 nM Daporinad (FK866, APO866) ++++ K:: 0.4 nM A922500 +++ IC ₅₀ : 7~24 nM	
Daporinad (FK866, APO866) ++++ K: 0.4 nM A922500 ++ ICo: 7~24 nM	
A922500 ++ ICso: 7~24 nM	
Lomeguatrib ++ IC ₅₀ : 5 nM	
FTI 277 HCI ++++ IC ₅₀ : 500 pM	
LB42708 +++ IC50: 0.8~2 nM	
PF-04620110 ++ IC50: 19 nM	
Tolcapone + Ki: 30 nM	

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

S2797 Lonafarnib (SCH66336)

Lonafarnib is an orally bioavailable FPTase inhibitor for H-ras, K-ras-4B and N-ras with IC50 of 1.9 nM. 5.2 nM and 2.8 nM in cell-free assays. respectively. Phase 3.

Size 5 mg 10 mg 10 mM/1 mL

HMG-CoA Reductase Inhibitors

Inhibitory Selectivity

Inhibitor Name	HMG-CoA Reductase
Simvastatin	++++ K _i : 0.1-0.2 nM
Rosuvastatin Calcium	++ IC ₅₀ : 11 nM
Lovastatin	+++ IC50: 3.4 nM
Fluvastatin Sodium	+++ IC50: 8 nM
Pravastatin sodium	++ IC ₅₀ : 5.6 μM
Clinofibrate	+ IC ₅₀ : 0.47 mM
Atorvastatin Calcium	1
Mevastatin	٧

Notes: 1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "V" refers to compounds which do inhibitory effects on the related isoform, but without specific value

S2169 Rosuvastatin Calcium (ZD4522)

Rosuvastatin Calcium is a competitive inhibitor of HMG-CoA reductase with IC50 of 11 nM in a cell-free assay. Size 50 mg 100 mg 1 g 10 mM/1 mL

S2061 Lovastatin (MK-803)

Lovastatin is an inhibitor of HMG-CoA reductase with IC50 of 3.4 nM, used for lowering cholesterol (hypolipidemic agent). Size 50 mg 200 mg 10 mM/1 mL



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S1909 Fluvastatin Sodium (XU-62-320)

,NH6	Fluvastatin Sodium inhibits HMG-CoA reductase activity nM in a cell-free assay.							of 8
0	Size	50 mg	5 g	10 mM/1 mL		d'	G**	-L _{ONs}



www.selleckchem.com





Size 50 mg

Metabolis

Gamma-secretase Inhibitors

Detailed product information is on page 92-93

S2077 Atorvastatin Calcium

Atorvastatin Calcium is an inhibitor of HMG-CoA reductase used as a cholesterol-lowering medication that blocks the production of cholesterol Of the

Size 50 mg 500 mg 10 mM/1 mL

Product Citation (1): BMC Pharmacol Toxicol, 2013, 14: 15 Data from [BMC Pharmacol Toxicol.

2013, 14: 151 Atorvastatin Calcium purchased from Selleck

S1759 Pitavastatin Calcium

Pitavastatin calcium, a novel member of the medication class of statins, is a calcium salt formulation of pitavastatin which is a highly effective HMG-CoA reductase inhibitor.

Size 10 mg 50 mg 200 mg

CETP Inhibitor

Inhibitory Selectivity

Inhibitor Name	CETP	
Anacetrapib (MK-0859)	+++ IC50: 7.9-11.8 nM	

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICses) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

S2748 Anacetrapib (МК-0859)

Anacetrapib (MK0859) is a potent, selective, reversible rhCETP and mutant CETP(C13S) inhibitor with IC50 of 7.9 nM and 11.8 nM, increases HDL-C and decreases LDL-C, does not increase aldosterone or blood pressure. Phase 3.

Size 5 mg 10 mg 10 mM/1 mL

Ferroptosis Inhibitors **Activators**

Ferroptosis Inhibitors

S7243 Ferrostatin-1 (Fer-1)

Ferrostatin-1 (Fer-1) is a potent and selective inhibitor of ferroptosis with EC50 of 60 nM. $O^{H} \dot{Q}_{a}$ Size 5 mg

S7699 Liproxstatin-1

Liproxstatin-1 is a potent ferroptosis inhibitor with an IC₅₀ of 22 nM. $\mathcal{O}_{n}^{\mathbb{I} \bigcap n}$ Size 5 mg 25 mg 100 mg

S7242 Erastin Erastin is a ferroptosis activator by acting on mitochondrial VDAC, mHo exhibiting selectivity for tumor cells bearing oncogenic RAS. Size 5 mg 50 mg S8155 RSL3 RSL3 is a ferroptosis activator in a VDAC-independent manner, exhibiting selectivity for tumor cells bearing oncogenic RAS. RSL3 binds, inactivates GPX4 and thus mediates GPX4-regulated ferroptosis. Size 5 mg 25 mg

Ferroptosis Activators

Calcitriol is a nonselective vitamin D receptor activator/agonist(VDRA),

exhibiting a 10-fold higher vitamin D receptor (VDR) binding

AhR Antagonists

Size 2 mg 5 mg 10 mM/1 mL

S2858 StemRegenin 1 (SR1)

StemRegenin 1 is an aryl hydrocarbon receptor (AhR) inhibitor with IC50 of 127 nM in a cell-free assay. Size 10 mg 50 mg 200 mg 10 mM/1 mL

S7711 CH-223191

Vitamin

S1466 Calcitriol

CH-223191 is a potent and specific aryl hydrocarbon receptor (AhR) antagonist with IC50 of 30 nM.

AhR Modulator

UM729 is an enhancer of aryl hydrocarbon receptor (AhR) antagonists. Size 5 mg 25 mg 100 mg



S7927 WZB117

WZB117 is an inhibitor of Glucose Transporter 1 (GLUT1). It inhibited cell proliferation in lung cancer A549 cells and breast cancer MCF7 cells with an IC50 of approximately 10 µM. Size 10 mg 50 mg 200 mg

Proteases



Proteasome Inhibitors

Detailed product information is on page 95-96

Caspase Inhibitors | Activator

Detailed product information is on page 57-58

HCV Protease Inhibitor

Inhibitory Selectivity

-	•
Inhibitor Name	HCV Protease
Daclatasvir (BMS-790052)	++++ EC50: 9~50 pM
Telaprevir (VX-950)	++ IC ₅₀ : 0.35 μM
Lomibuvir (VX-222, VCH-222)	+ ICso: 0.94~1.2 μM
Danoprevir (ITMN-191)	+++ IC50: 0.2~3.5 nM

Notes:

1. For more details, such as half maximal inhibitory concentrations (IC50s) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

Daclatasvir (BMS-790052) is a highly selective inhibitor of HCV NS5A with EC50 of 9-50 pM, for a broad range of HCV replicon genotypes and

No inhibitor treatment





S1482 Daclatasvir (BMS-790052, EBP883)

Product Citations (14): Nature, 2013, 501(7466): 237-41 Hepatology, 2014, 10.1002/hep.27197

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Data from [Antimicrob Agents Chemother, 2014, 58(1): 386-961 Daclatasvir purchased from Selleck

122

AhR Antagonists | Modulator

affinity(IC50=0.4 nM) than the selective VDRA paricalcitol.

Size 10 mg 50 mg 200 mg

S7510 UM729





DPP-4 Inhibitors

Inhibitory Selectivity

Inhibitor Name	DPP-4
Sitagliptin phosphate monohydrate	++ ICso: 19 nM
Linagliptin	++++ IC ₅₀ : 1 nM
Vildagliptin (LAF-237)	+++ IC50: 2.3 nM
Saxagliptin	+ ICso: 26 nM
Alogliptin	+++ IC50: <10 nM
Trelagliptin	1

Notes:

1. For more details, such as half maximal inhibitory concentrations (IC50S) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation.

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

S4002 Sitagliptin phosphate monohydrate (MK-0431)

Sitagliptin phosphate monohydrate is a potent inhibitor of DPP-IV with IC50 of 19 nM in Caco-2 cell extracts

Size 200 mg 10 mM/1 mL

S3031 Linagliptin (BI-1356)

Linagliptin is a highly potent, selective DPP-4 inhibitor with IC50 of 1 nM and exhibits a 10,000-fold higher selectivity for DPP-4 than for other dipeptidyl peptidases such as DPP-2, DPP-8, and DPP-9.

Size 5 mg 10 mg 10 mM/1 mL

S3033 Vildagliptin (LAF-237)

Vildagliptin (LAF-237) inhibits DPP-4 with IC50 of 2.3 nM. Size 10 mg 25 mg 10 mM/1 mL S. L.L



S1380 Lopinavir (ABT-378)

S1185 Ritonavir (ABT-538, A 84538)

Lopinavir is a potent HIV protease inhibitor with K of 1.3 pM in a cell-free assav



Atazanavir Sulfate is a HIV protease inhibitor with Ki of 2.66 nM in a cell-free assay Size 5 mg 10 mg 50 mg 10 mM/1 mL

HIV Protease Inhibitors

Inhibitory Selectivity

Inhibitor Name	HIV Protease	Other
Lopinavir	++++ K _i : 1.3 pM	
Atazanavir Sulfate	++ Ki: 2.66 nM	
Amprenavir	+ IC50: 14.6 ng/mL	
Nelfinavir Mesylate	+++ Ki: 2 nM	
Ritonavir	√	CYP3A4
Darunavir Ethanolate	*	
Limonin	1	

Notes:

1. For more details, such as half maximal inhibitory concentrations (IC50S) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "\" refers to compounds which do inhibitory effects on the related isoform, but without S7155 Batimastat (BB-94) specific value

MMP Inhibitors

Inhibitory \$	Selectivity
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Inhibitor Name	MMP
Batimastat (BB-94)	+++ ICso: 3~20 nM
llomastat (GM6001, Galardin)	++++ Ki: 0.1~3.7 nM
SB-3CT	+ K _i : 13.9~600 nM
Marimastat (BB-2516)	+++ ICso: 3~230 nM
NSC 405020	V
Nobiletin	√
Notes:	

1. For more details, such as half maximal inhibitory concentrations (IC50s) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

Size 1 mg 10 mg

Batimastat (BB-94) is a potent, broad spectrum matrix metalloprotease (MMP) inhibitor for MMP-1, MMP-2, MMP-9, MMP-7 and MMP-3 with IC50 of 3 nM, 4 nM, 4 nM, 6 nM and 20 nM, respectively. Also inhibits the activitity of other metalloproteases, such as ADAM17.



S7380 Leupeptin Hemisulfate

Leupeptin Hemisulfate is a reversible inhibitor of serine and cysteine proteases. It inhibits cathepsin B (Ki = 6 nM), calpain (Ki = 10 nM), trypsin (K_i = 35 nM), plasmin (K_i = 3.4μ M), and kallikrein (K_i = 19μ M), and has no effect against chymotrypsin, elastase, renin, or pepsin.

Size 10 mg 50 mg

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Size

S7396 Calpeptin

Calpeptin is a potent, cell-permeable calpain inhibitor with ID50 of 52 nM, 34 nM, 138 nM, and 40 nM for Calpain I (porcine erythrocytes), Calpain II (porcine kidney), Papainb, and Calpain I (human platelets), respectively.

10 mg 50 mg 200 mg

S4163 Doxycycline Hyclate

Size 5 mg 25 mg 100 mg

S7157 Ilomastat (GM6001, Galardin)

respectively

Size 5 mg

S7430 SB-3CT

Doxycycline is a member of the tetracycline antibiotics group, and is commonly used to treat a variety of infections. It is also an inhibitor of matrix metallo-proteinases (MMP) Size 50 mg

llomastat (GM6001, Galardin) is a broad spectrum matrix

metalloprotease (MMP) inhibitor for MMP-1, MMP-2, MMP-3, MMP-7,

MMP-8, MMP-9, MMP-12, MMP-14, and MMP-26 with K of 0.4 nM, 0.5

nM, 27 nM, 3.7 nM, 0.1 nM, 0.2 nM, 3.6 nM, 13.4 nM, 0.36 nM,

SB-3CT is an effective and selective gelatinase inhibitor with K of 13.9

nM and 600 nM for MMP-2 and MMP-9, respectively.

S7420 CA-074 methyl ester (CA-074 Me) new

CA-074 Me is a membrane-permeable derivative of CA-074 and acts as an irreversible cathepsin B inhibitor. 5 mg 25 mg

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Cysteine Protease Inhibitors

Inhibitory Selectivity

Inhibitor Name	Cysteine Protease	Other
Odanacatib (MK-0822)	++++ IC ₅₀ : 0.2 nM	
E-64	+++ IC50: 9 nM	
PD 151746	+ IC50: 5.33 μM	
Calpeptin	++ ID ₅₀ : 52 nM	
Cathepsin Inhibitor 1	+++ pIC ₅₀ : 5.2	
PMSF	√	chymotrypsin
Aloxistatin	√	
Loxistatin Acid (E-64C)	√	
Leupeptin Hemisulfate	√	serine protease
Z-FA-FMK	√	
MG-101 (ALLN)	1	

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.con 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation

3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without snecific value

S7379 E-64

E-64 is an irreversible and selective cysteine protease inhibitor, and also inhibits papain, calpain, and cathepsins B and H, but not serine proteases or aspartic proteases. The IC₅₀ for papain is 9 nM.

Size 10 mg 25 mg

S7393 Aloxistatin (E-64d)

Aloxistatin is an irreversible and membrane-permeable cysteine protease inhibitor with blood platelet aggregation inhibiting activity. Size 2 mg 5 mg

S7386 MG-101 (ALLN)

MG-101 (ALLN) is a cell-permeable and potent inhibitor of cysteine proteases including calpains and lysosomal cathepsins

Serine Protease Inhibitors

Inhibitory Selectivity

Inhibitor Name	Serine Protease	Other
Gabexate Mesylate	++ IC50: 0.19 μM	
Aprotinin	+++ K _i : 9.5 nM	Thrombin, Trypsin, kallikrein
Alvelestat (AZD9668)	++++ IC50: 12 nM	
Nafamostat Mesylate	1	
PMSF	√	cysteine protease
Sivelestat (ONO-5046)	1	
Leupeptin Hemisulfate	1	Cysteine protease
AEBSF HCI	1	

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "V" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

S7378 AEBSF HCI

AEBSF HCI is a broad spectrum, irreversible serine protease inhibitor. Size 100 mg 250 mg 500 mg

S7218 Alvelestat (AZD9668)

Alvelestat (AZD9668) is an oral, highly selective inhibitor of neutrophil elastase (NE) with IC50 and Ki of 12 nM and 9.4 nM, at least 600-fold more selective over other serine proteases. Phase 2. Size 5 mg 25 mg

Sivelestat is a potent and selective inhibitor of neutrophil elastase with IC50 of 44nM. It almost shows no activity at a range of other proteases. Size 10 mg 50 mg 200 mg

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S8136 Sivelestat (ONO-5046)

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Immunology & Inflammation



PD-1/PD-L1 Inhibitors

S7912 BMS202 (PD-1/PD-L1 inhibitor 2)

BMS202 (PD-1/PD-L1 inhibitor 2) is a small-molecule PD-1/PD-L1 PD-1/PD-L1 inhibitor 1 is a small-molecule inhibitor of PD-1/PD-L1 interaction inhibitor with IC50 of 18 nM. Size 5 mg 25 mg QL.O.Y

S8158 PD-1/PD-L1 Inhibitor 3

PD-1/PD-L1 Inhibitor 3 (Programmed Death-1/Programmed Death -Ligand 1 Inhibitor 3) is a Macrocyclic inhibitor of PD-1/PD-L1 interaction with IC50 of 5.6 nM. Size 1 mg 5 mg





S7911 PD-1/PD-L1 inhibitor 1

Size 5 mg 25 mg

AUNP-12, a new immune checkpoint modulator, is an inhibitor of the PD-1 pathway





S8133 Resiguimod

GKT137831 is a potent, dual NADPH oxidase NOX1/NOX4 inhibitor

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S3013 Plerixafor 8HCI (AMD3100 8HCI)

Size 5 mg 10 mg 50 mg

S8030 Plerixafor (AMD3100)

cell-free assays, respectively.

Size 5 mg 10 mg 50 mg

over the other 7-TMRs tested.

Size 10 mg 50 mg 200 mg

S1322 Dexamethasone (DHAP)

SB225002 is a potent, and selective CXCR2 antagonist with IC $_{50}$ of 22 nM for inhibiting interleukin IL-8 binding to CXCR2, > 150-fold selectivity

Celecoxib is a selective COX-2 inhibitor with IC50 of 40 nM in Sf9 cells

Plerixafor 8HCI (AMD3100 8HCI) is the hydrochloride of Plerixafor, a

chemokine receptor antagonist for CXCR4 and CXCL12-mediated

chemotaxis with IC50 of 44 nM and 5.7 nM in cell-free assays,

Cyclosporin A is an immunosuppressive agent, binds to the cyclophilin

and then inhibits calcineurin with IC₅₀ of 7 nM in a cell-free assay, widely

used in organ transplantation to prevent rejection.

S7651 SB225002

S1261 Celecoxib

Size 100 mg 1 g

respectively.

Size 50 mg 10 mM/1 mL

S2286 Cyclosporin A

Size 50 mg 5 g 10 mM/1 mL

respectively.

with K_i of 110 nM and 140 nM, respectivelyl; ~10-fold selectivity towards NOX1, 4 and 5 over NOX2, does not inhibit XO or scavange ROS/RNS.

Size 5 mg 25 mg 100 mg

S2003 Maraviroc

Maraviroc is a CCR5 antagonist for MIP-1a, MIP-1B and RANTES with IC50 of 3.3 nM, 7.2 nM and 5.2 nM in cell-free assays, respectively.

Size 5 mg 25 mg 100 mg 10 mM/1 mL

S1623 Acetylcysteine

Acetylcysteine(N-acetyl-I-cysteine) is a ROS(reactive oxygen species) inhibitor that antagonizes the activity of proteasome inhibitors. It is also a tumor necrosis factor production inhibitor, used mainly as a mucolytic, protects against acetaminophen overdose-induced hepatotoxicity by maintaining or restoring hepatic concentrations of glutathione.

Size 10 mg 50 mg 10 mM/1 mL

S1848 Curcumin

Curcumin is the principal curcuminoid of the popular Indian spice turmeric, which is a member of the ginger family (Zingiberaceae). It is an inhibitor of p300 histone acetylatransferase (IC $_{50}\text{--}25~\mu\text{M})$ and Histone deacetylase; activates Nrf2 pathway and supresses the activation of transcription factor NF-KB.

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S8549 AUNP-12

Microbiology



HCV Protease Inhibitor

Detailed product information is on page 123

S2005 Raltegravir (MK-0518)

Size

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Inhibitory Selectivity

Integrase Inhibitors

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Inhibitor Name	Integrase
Raltegravir (MK-0518)	+ IC50: 40-90 nM
Elvitegravir (GS-9137, JTK-303)	++++ ICso: 0.7-2.8 nM
Dolutegravir (GSK1349572)	+++ IC50: 2.7 nM
BMS-707035	++ ICso: 15 nM
MK-2048	+++ ICso: 1.5-2.6 nM
Dolutegravir Sodium	+++ ICso: 2.7 nM
Cabotegravir (GSK744, GSK1265744)	1

Notes:

1. For more details, such as half maximal inhibitory concentrations (ICsos) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com.

2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value.

Raltegravir (MK-0518) is a potent integrase (IN) inhibitor for WT and S217Q PFV IN with IC50 of 90 nM and 40 nM in cell-free assays, respectively. It shows greater than 1000-fold selectivity for HIV-1 IN over several related Mg2+-dependent enzyme such as HCV polymerase, HIV reverse transcriptase, HIV RNaseH and human α-, β-, γ-polymerases.

HIV Protease Inhibitors

Detailed product information is on page 124





Elvitegravir (GS-9137, JTK-303) is an HIV integrase inhibitor for HIV-1 IIIB, HIV-2 EHO and HIV-2 ROD with IC50 of 0.7 nM, 2.8 nM and 1.4 nM, respectively

Size 10 mg 50 mg 10 mM/1 mL



S2667 Dolutegravir (GSK1349572)

Dolutegravir (GSK1349572) is a two-metal-binding HIV integrase inhibitor with IC50 of 2.7 nM, modest activity against raltegravir-resistant signature mutants Y143R, Q148K, N155H, and G140S/Q148H.



S4642 Dolutegravir Sodium

Dolutegravir is a HIV integrase inhibitor with IC50 of 2.7 nM. Size 5 mg 25 mg

Reverse Transcriptase Inhibitors

Inhibitory Selectivity

Inhibitor Name	Ir	ntegrase	Other
Didanosine	++	IC50: 490 nM	
Dapivirine (TMC120)	+++	IC50: 24 nM	
Tenofovir	1		
Tenofovir Disoproxil Fumarate	1		
Emtricitabine	1		
Entecavir Hydrate	1		
Adefovir Dipivoxil	1		
Nevirapine	1		
Lamivudine	1		
Stavudine (d4T)	1		
Telbivudine	1		
Etravirine (TMC125)	1		

Integrase / Reverse Transcriptase

Inhibitory Selectivity

Inhibitor Name	Integrase	Other
Zidovudine	1	
Zalcitabine	1	
Abacavir sulfate	1	
Foscarnet Sodium	1	RNA polymerase, DNA polymerase
Rilpivirine	1	
Salicylanilide	1	integrase

Sa Notes:

1. For more details, such as half maximal inhibitory concentrations (IC505) and working concentrations of each inhibitor, please visit the website of www.selleckchem.com. 2. "+" indicates inhibitory effect. Increased inhibition is marked by a higher "+" designation 3. Red "v" refers to compounds which do inhibitory effects on the related isoform, but without specific value

S1401 Tenofovir

Tenofovir blocks reverse transcriptase and hepatitis B virus infections. Size 5 mg 20 mg 50 mg 10 mM/1 mL



Tenofovir Disoproxil Fumarate belongs to a class of antiretroviral drugs, it inhibits the activity of HIV reverse transcriptase by competing with the natural substrate deoxyadenosine 5'-triphosphate and, afte incorporation into DNA, by DNA chain termination.

Size 10 mg 50 mg 10 mM/1 mL

S1704 Emtricitabine

Emtricitabine (FTC) is a new nucleoside agent that has activity against both human immunodeficiency virus (HIV) and hepatitis B virus. It is a reverse transcriptase inhibitor. Intracellular half-life is 39 h. Size 10 mg 50 mg 200 mg 10 mM/1 mL J.C.

Nevirapine is a non-nucleoside reverse transcriptase inhibitor (NNRTI) used to treat HIV-1 infection and AIDS. Size 5 mg 25 mg 100 mg 10 mM/1 mL d d

S1706 Lamivudine

Lamivudine is a potent nucleoside analog reverse transcriptase inhibitor, used for treatment of chronic HBV and HIV/AIDS. It works by blocking the HIV reverse transcriptase and hepatitis B virus polymerase. 1° Size 10 mg 25 mg 50 mg 10 mM/1 mL

S2579 Zidovudine

Zidovudine is a nucleoside analogue reverse transcripta	se inhibitor,
Size 25 mg 100 mg 1g	Ĵ
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new



CCR / Antifection

CCR Antagonist

S2003 Maraviroc (UK-427857)

Maraviroc is a CCR5 antagonist for MIP-1g. MIP-1g and RANTES with IC50 of 3.3 nM, 7.2 nM and 5.2 nM in cell-free assays, respectively.

Size 5 mg 25 mg 100 mg 10 mM/1 mL



Product Citations (5): Cancer Res, 2014, 74(23): 7103-14 J Neuroimmune Pharmacol, 2014, 9(5):

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Size 250 mg

viruses, used to prevent influenza.

Oseltamivir Phosphate

S2908 Hygromycin B

S2597

Size 1 g

inhibitor.

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Hygromycin B, a selective antibiotic that is effective on most bacteria, fungi and higher eukaryotes, inhibits protein synthesis by interfering with translocation and causing mistranslation at the 70S ribosome. Size 250 mg

Oseltamivir Phosphate is a potent and selective inhibitor of the neuraminidase that is essential for replication of influenza A and B



S3028 Geneticin (G418 Sulfate)

S3073 Caspofungin Acetate

S3162 Tylosin tartrate

Size 50 mg 10 mM/1 mL

Size 50 mg

Geneticin (G418 Sulfate), an aminoglycoside antibiotic, is an elongation inhibitor of 80 S ribosomes that blocks polypeptide synthesis by inhibiting the elongation step in both prokaryotic and eukaryotic cells.



Antifection

S1517 Natamycin

Natamycin, a natural and versatile anti-fungal agent during fermentation by the bacterium Streptomyces natalensis, commonly found in soil; with little to no flavour interference.

Size 50 mg 100 mg 200 mg 10 mM/1 mL

S1878 Ganciclovir

Ganciclovir is an antiviral drug for feline herpesvirus type-1 with IC50 of 5.2 µM in a cell-free assay.

Size 50 mg 250 mg 10 mM/1 mL

S2265 Artesunate

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Artesunate is a part of the artemisinin group of agents with an IC50 of < 5 µM for small cell lung carcinoma cell line H69. It is a potential inhibitor of STAT-3 and exhibits selective cytotoxicity of cancer cells over normal cells in vitro; A potent inhibitor of EXP1.

Size 10 mg 50 mg 200 mg 10 mM/1 mL



Tylosin tartrate is a macrolide antibiotic approved for the control of mycoplasmosis in poultry.

S7417 Puromycin 2HCI

Puromycin 2HCl is an aminonucleoside antibiotic, which acts as a protein synthesis inhibitor.



Phosphorylase / IL Receptor / Thrombin / Liver X Receptor / PKA / Substance P

Phosphorylase Inhibitor

S2717 CP-91149 Licensed by Pfizer

CP-91149 is a selective glycogen phosphorylase (GP) inhibitor with IC50 of 0.13 μM in the presence of glucose, 5- to 10-fold less potent in the absence of glucose. Size 5 mg 10 mg 100 mg 10 mM/1 mL

Liver X Receptor Agonists

S2630 GW3965 HCI

S7076 T0901317

GW3965 HCl is a potent, selective LXR agonist for hLXRa and hLXRβ with EC50 of 190 and 30 nM in cell-free assays, respectively.









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T0901317 is a potent and selective agonist for both LXR and FXR, with EC50 of ~50 nM and 5 µM, respectively. Size 25 mg 100 mg 0°°O-E

IL Receptor Inhibitor | Modulator

IL Receptor Inhibitor

S4028 Dexamethasone Sodium Phosphate

Dexamethasone Sodium Phosphate is a potent synthetic member of the glucocorticoid class of steroid drugs, and an interleukin receptor modulator that has anti-inflammatory and immunosuppressant effects. Size 50 mg



IL Receptor Modulator

S1322 Dexamethasone (DHAP)

Dexamethasone (DHAP) is a potent synthetic member of the glucocorticoid class of steroid drugs, and aninterleukin receptor modulator that has anti-inflammatory and immunosuppressant effects.

Size 50 mg 10 mM/1 mL



Thrombin Inhibitor

S2196 Dabigatran (BIBR 953)

Size 5 mg 10 mg 50 mg

IC50 of 9.3 nM in a cell-free assay.

PKA Inhibitor | Activators

PKA Inhibitor

S1582 H 89 2HCI

H 89 2HCl is a potent PKA inhibitor with Ki of 48 nM in a cell-free assay, 10-fold selective for PKA than PKG, 500-fold greater selectivity than PKC, MLCK, calmodulin kinase II and casein kinase I/II. $({\rm Span}^{\rm res})^{\rm res}$



PKA Activators

S7857 8-Bromo-cAMP

8-bromo-cAMP is a cell permeable analog of cAMP that activates cyclic-AMP-dependent protein kinase with a Ka value of 0.05 µM; and a PKA activator.



S7858 Dibutyryl-cAMP (Bucladesine)

Dibutyryl-cAMP (Bucladesine) is a cell-permeable PKA activator by mimicing the action of endogenous cAMP.

Size 100 mg 500 mg

Substance P Antagonist

S1189 Aprepitant (MK-0869, L-754030)

Aprepitant is a potent and selective neurokinin-1 receptor antagonist with IC₅₀ of 0.1 nM.

Size 2 mg 10 mg 25 mg 10 mM/1 mL



Dabigatran (BIBR 953) is a potent nonpeptide thrombin inhibitor with an $= \frac{1}{\sqrt{2}} \frac{1}{\sqrt{$



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No Cod

Others

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FXR / gp120/CD4 / Phosphatase / NADPH Oxidase / PTEN / Others

FXR Agonists

S2782 GW4064

GW4064 is an agonist of farnesoid X receptor (FXR) with EC₅₀ of 65 nM Size 5 mg 25 mg 100 mg in CV1 cell line and displays no activity at other nuclear receptors at concentrations up to 1 µM

Size 5 mg 25 mg 50 mg 10 mM/1 mL

S2694 Turofexorate Isopropyl (XL335, Fxr 450)

Turofexorate Isopropyl (XL335) is a potent, selective FXR agonist with EC50 of 4 nM, highly selective versus other nuclear receptors, such as LXR. PPAR. ER and etc. Phase 1.

Size 5 mg 10 mg 50 mg 10 mM/1 mL

S7660 Obeticholic Acid

Obeticholic Acid is a potent and selective farnesoid X receptor (FXR) agonist with EC50 of 99 nM. Phase 3. Size 5 mg 25 mg 100 mg

Others

gp120/CD4 Inhibitor

S2632 BMS-378806

S1949 Menadione

as a nutritional supplement.

Size 50 mg 10 mM/1 mL

BMS-378806 selectively inhibits the binding of HIV-1 gp120 to the CD4 receptor with EC50 of 0.85-26.5 nM in virus. 9-0

Size 5 mg 10 mg 50 mg 10 mM/1 mL

S7171 GKT137831 GKT137831 is a potent, dual NADPH oxidase NOX1/NOX4 inhibitor

with K_i of 110 nM and 140 nM, respectivelyl; ~10-fold selectivity towards NOX1, 4 and 5 over NOX2, does not inhibit XO or scavange ROS/RNS



Others

S5003 Tacrolimus (FK506)

Tacrolimus (FK506) is a 23-membered macrolide lactone, it reduces peptidyl-prolyl isomerase activity in T cells by binding to the immunophilin FKBP12 (FK506 binding protein) creating a new complex. Size 50 mg 100 mg 500 mg 10 mM/1 mL

S1212 Bendamustine HCI

Bendamustine HCI is a DNA-damaging agent with IC $_{50}$ of 50 μM in cell-free assay. Size 25 mg 100 mg 10 mM/1 mL

S1290 Celastrol

Celastrol is a potent proteasome inhibitor for the chymotrypsin-like activity of a purified 20S proteasome with IC50 of 2.5 µM. Size 10 mg 50 mg 100 mg

S1373 Daptomycin

Daptomycin is a novel antibiotic with rapid in vitro bactericidal activity against gram-positive organisms. Size 20 mg 50 mg 100 mg

S8278 SHP099 dihydrochloride

Phosphatase Inhibitors

SHP099 is a highly potent, selective and orally bioavailable smallmolecule SHP2 inhibitor with an IC $_{\rm 50}$ value of 0.071 μM and shows no activity against SHP1

NADPH Oxidase Inhibitors

Apocynin is a selective NADPH-oxidase inhibitor with IC50 of 10 µM.

Menadione(Vitamin K3), a fat-soluble compound, is an inhibitor of

Cdc25 phosphatase and mitochondrial DNA polymerase y (pol y), used

Size 5 mg 25 mg 100 mg

S2485 Mitoxantrone HCI Mitoxantrone is a type II topoisomerase inhibitor with IC50 of 2.0 µM, 0.42 mM for HepG2 and MCF-7/wt cells, respectively. $\sum_{n=1}^{n} \sum_{n \in \mathbb{N}}$ 50 mg 100 mg 10 mM/1 mL Size

S1680 Disulfiram

Disulfiram is a specific inhibitor of aldehyde-dehydrogenase (ALDH1), used for the treatment of chronic alcoholism by producing an acute sensitivity to alcohol 50 mg 10 mM/1 mL

S1692 Busulfan

Busulfan is a cell cycle non-specific alkylating antineoplastic agent. Size 50 mg 10 mM/1 mL

S1709 Estradiol

Estradiol, or more precisely, 17β-estradiol, is a human sex hormone and steroid, and the primary female sex hormone. Size 50 mg 10 mM/1 mL

S1653 Tretinoin

Tretinoin, which is a ligand for both the retinoic acid receptor (RAR) and the retinoid X receptor (RXR), can induce granulocytic differentiation and apoptosis in acute promyelocytic leukemia (APL) cells. Size 50 mg 10 mM/1 mL

S1896 Hydroxyurea

Hydroxyurea is an antineoplastic agent that inhibits DNA synthesis through the inhibition of ribonucleoside diphosphate reductase. Size 10 mg 50 mg 200 mg 10 mM/1 mL

S1950 Metformin HCI

Metformin HCI decreases hyperglycemia in hepatocytes primarily by suppressing glucose production by the liver (hepatic gluconeogenesis). Size 50 mg 5 g 10 mM/1 mL

S1899 Nicotinamide (Vitamin B3)

Nicotinamide (Vitamin B3), a water-soluble vitamin, is an active component of coenzymes NAD and NADP, and also act as an inhibitor of sirtuins. Size 50 mg 10 mM/1 mL

S1792 Simvastatin

Simvastatin is a competitive inhibitor of HMG-CoA reductase with Ki of 0.1-0.2 nM in cell-free assays. Size 25 mg 100 mg

S2286 Cyclosporin A

Cyclosporin A is an immunosuppressive agent, binds to the cyclophilin and then inhibits calcineurin with IC50 of 7 nM in a cell-free assay, widely used in organ transplantation to prevent rejection. Size 50 mg 5 g 10 mM/1 mL

S1786 Verteporfin

Verteporfin is a potent second-generation photosensitizing agent derived from porphyrin in endothelial cel. Size 10 mg 50 mg

S2476 Itraconazole

Itraconazole is a relatively potent inhibitor of CYP3A4 with IC50 of 6.1 nM, used as a triazole antifungal agent. Size 100 mg 200 mg

S1696 Hydrocortisone

Hydrocortisone is a steroid hormone or glucocorticoid produced by the adrenal gland. Size 50 mg 10 mM/1 mL

S2590 Pioglitazone

Pioglitazone is a selective peroxisome proliferator-activated receptor-gamma (PPARy) agonist, used to treat diabetes; A weak activator for full-length hPPARa, but not full-length hPPARo. Size 10 mg 50 mg 200 mg 10 mM/1 mL

S2057 Cyclophosphamide Monohydrate

Cyclophosphamide Monohydrate is a nitrogen mustard alkylating agent, it attaches the alkyl group to the guanine base of DNA, shown to crosslink DNA, causing strand breakage and inducing mutations. Size 50 mg 5 g

S2858 StemRegenin 1 (SR1)

StemRegenin 1 is an aryl hydrocarbon receptor (AhR) inhibitor with IC50 of 127 nM in a cell-free assay

Size 10 mg 100 mg 200 mg 10 mM/1 mL

S3022 Cabazitaxel

S2877 L-NAME HCI

Cabazitaxel is a semi-synthetic derivative of a natural taxoid that kills cancer cells by inhibiting cell division and growth. Cabazitaxel exerts its effects by inhibiting microtubule growth and assembly, processes that are essential for cells to divide.

Size 5 mg 10 mg 10 mM/1 mL

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L-NAME HCl is a nonselective inhibitor of nitric oxide synthetases (NOS) for nNOS (bovine), eNOS (human), and iNOS (murine), with Ki of 15 nM, 39 nM and 4.4 µM, respectively. Size 100 mg

S3190 N6-methyladenosine (m6A)

N6-methyladenosine (m6A) is a base modified analog of adenosine and is found as a minor nucleoside in natural RNAs. Size 50 mg

S4202 Verapamil HCI

Verapamil HCl is an L-type calcium channel blocker that is a class IV anti-arrhythmia agent. Size 50 mg

S4227 Fidaxomicin

Fidaxomicin is a narrow spectrum macrocyclic antibiotic that inhibits RNA polymerase sigma subunit. Size 50 mg

S7272 4µ8C

4u8C is a potent and selective IRE1 Rnase inhibitor with IC50 of 76 nM. Size 10 mg 50 mg

S7534 BAPTA-AM

BAPTA-AM is a selective, membrane-permeable calcium chelator. Size 10 mg 50 mg

S7381 Pepstatin A

Pepstatin A is a potent aspartic protease inhibitor, and also inhibits HIV replication Size 10 mg 50 mg 200 mg

S7209 GSK650394

GSK650394 is a serum- and glucocorticoid-regulated kinase-1 inhibitor with IC50 of 62 nM and 103 nM for SGK1 and SGK2, respectively. Size 5 mg 25 mg 100 mg

Size 1 g

S2425 Apocynin

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37537 LB-100	S7809 MCC950 (CP-456773)
B-100 is a water soluble protein phosphatase 2A (PP2A) inhibitor with C_{50} s of 0.85 µM and 3.87 µM in BxPc-3 and Panc-1 cells.	MCC950 sodium salt is a potent, selective inhibitor of NLRP3 with IC ₅₀ of 7.5 nM in BMDMs; but not the AIM2, NLRC4 or NLRP1 inflammasomes. <u>Size 10 mg 50 mg 200 mg</u>
37655 CB-839	57020 A7D0055

CB-839 is a potent, selective, and orally bioavailable glutaminase inhibitor with IC50 of 24 nM for recombinant human GAC. Phase 1. Size 5 mg 25 mg 100 mg

transporter 1 (MCT1) inhibitor with a binding affinity of 1.6 nM, 6-fold selective over MCT2. Phase 1. Size 5 mg 25 mg

S7753 BPTES

BPTES is a potent and selective Glutaminase GLS1 (KGA) inhibitor with IC_{\rm 50} of 0.16 $\mu M.$ It has no effect on glutamate dehydrogenase activity and causes only a very slight inhibition of y-glutamyl transpeptidase activity. Size 10 mg

S8368 LM10 new LM10 is a selective tryptophan 2,3-dioxygenase (TDO) inhibitor with IC50 values of 0.62 and 2 µM for human and mouse TDO, respectively.

AZD3965 is a potent, selective and orally available monocarboxylate

Size 10 mg 50 mg 200 mg

S7771 STF-083010

STF-083010 is a specific IRE1 α endonuclease inhibitor without affecting its kinase activity. Size 10 mg 50 mg 200 mg

Others