

Apoptosis

Apoptosis

Apoptosis is a distinctive form of cell death exhibiting specific morphological and biochemical characteristics, including cell membrane blebbing, chromatin condensation, genomic DNA fragmentation, and exposure of specific phagocytosis signaling molecules on the cell surface. Cells undergoing apoptosis differ from those dying through necrosis. Necrotic cells are usually recognized by the immune system as a danger signal and, thus, resulting in inflammation; in contrast, apoptotic death is quiet and orderly.

There are two major pathways of apoptotic cell death induction: The intrinsic pathway, also called the Bcl-2-regulated or mitochondrial pathway, is activated by various developmental cues or cytotoxic insults, such as viral infection, DNA damage and growth-factor deprivation, and is strictly controlled by the BCL-2 family of proteins. The extrinsic or death-receptor pathway is triggered by ligation of death receptors (members of the tumor necrosis factor (TNF) receptor family, such as Fas or TNF receptor-1 (TNFR1)) that contain an intracellular death domain, which can recruit and activate caspase-8 through the adaptor protein Fas-associated death domain (FADD; also known as MORT1) at the cell surface. This recruitment causes subsequent activation of downstream (effector) caspases, such as caspase-3, -6 or -7, without any involvement of the BCL-2 family.

Studies suggest that alterations in cell survival contribute to the pathogenesis of a number of human diseases, including cancer, viral infections, autoimmune diseases, neurodegenerative disorders, and AIDS (acquired immunodeficiency syndrome). Treatments designed to specifically alter the apoptotic threshold may have the potential to change the natural progression of some of these diseases.

Apoptosis Inhibitors, Antagonists, Activators, Modulators & Inducers





(R)-Verapamil D/ hydrochloride		(R)-Verapamil hydrochloride	
((R)-(+)-Verapamil D7 hydrochloride) (R)-Verapamil D7 hydrochloride ((R)-(+)-Verapamil	Cat. No.: HY-135336S	((R)-(+)-Verapamil hydrochloride) (R)-Verapamil hydrochloride ((R)-(+)-Verapamil	Cat. No.: HY-135336
D7 hydrochloride) is a deuterium labeled (R)-Verapamil bydrochloride (R)-Verapamil		hydrochloride) is a P-Glycoprotein inhibitor. (R)-Verapamil hydrochloride blocks MRP1 mediated	I den
hydrochloride ((R)-(+)-Verapamil hydrochloride) is	N N N N	transport, resulting in chemosensitization of	- Q ~ N ~ D
a P-Glycoprotein inhibitor.	H-CI _O _O	MRP1-overexpressing cells to anticancer drugs.	нсі — ⁰ —
Purity: >98%	Const Contract Toront (1997)	Purity: 98.54%	
Clinical Data: No Development Reported		Clinical Data: Launched Size: 10 mM x 1 mL 5 mg 10 mg 50 mg 100 mg	
Jize. Ting		512e. 10 million × 1 millio, 5 millio, 50 millio, 100 millio	
(R)-CR8		(Rac)-Antineoplaston A10	
(CR8, (R)-Isomer)	Cat. No.: HY-18340		Cat. No.: HY-128553A
(R)-CR8 (CR8), a second-generation analog of Roscovitine, is a potent CDK1/2/5/7/9 inhibitor.	HOTHIN	(rac)-Antineoplaston A10 is the racemate of Antineoplaston A10. Antineoplaston A10 is a Ras inhibitor potentially for the treatment of glioma, lymphoma, astrocytoma and breast cancer.	
D :: 00.000/	N	D :: 000/	н
Clinical Data: No Development Reported	\bigtriangledown	Clinical Data: No Development Reported	
Size: 10 mM × 1 mL, 5 mg, 10 mg		Size: 1 mg, 5 mg	
(Rac)-Hesperetin	Cat. No. LIV. NO1694	(Rac)-Hesperetin-13C,d3	Cat No. LIV NO169461
			Cat. NO.: HT-NU106AS1
(Rac)-Hesperetin is the racemate of Hesperetin. Hesperetin is a natural flavanone, and acts as a		(Rac)-Hesperetin-13C,d3 is the 13C- and deuterium labeled. (Rac)-Hesperetin is the racemate of	.727
potent and broad-spectrum inhibitor against human		Hesperetin. Hesperetin is a natural flavanone, and acts as a potent and broad-spectrum inhibitor	HQ ~ Q
p38 MAPK activation.	HOLOGIA	against human UGT activity. Hesperetin induces	TTT OH
Durity: 08.20%	~ 0	apoptosis via p38 MAPK activation.	OH O
Clinical Data: No Development Reported		Clinical Data: No Development Reported	
Size: 100 mg		Size: 1 mg, 5 mg	
(Dac) Llogramatin d2		(Dec) Idversieleis esid	
(Kac)-Hesperetin-d3	Cat No HY-N0168AS	(Rac)-IOTOXIOTEIC ACIO (2-Hvdroxvoleic acid: 2-OHOA)	Cat No : HY-129467
(Bac)-Hesperetin-d3 is the deuterium labeled		(Rac)-Idrovioleic acid (2-Hydroxyoleic acid) is a	
(Rac)-Hesperetin. (Rac)-Hesperetin is the racemate	04.0	synthetic oleic acid (OA) derivative that binds to	
of Hesperetin. Hesperetin is a natural flavanone, and acts as a potent and broad-spectrum inhibitor	Ϋ́́,	the plasma membrane and alters lipid organization. (Rac)-Idroxioleic acid has anti-tumor effect.	l
against human UGT activity. Hesperetin induces	HOTOTOT		он от от от от от он _{он}
Purity: >98%		Purity: 96.49%	
Clinical Data: No Development Reported		Clinical Data: Phase 2	
Size: 1 mg, 10 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg	
(KaC)-IOFOXIOIEIC aCIO-OL/ (2-Hydroxyoleic acid-d17: 2-OHOA-d17)	Cat No: HV-1294675	(KaC)-INGOXIMOG (1-Methyl-DL-tryptophan: (Rac)-NI G-8189)	Cat No: HV-133807
(Pac)-Idrovioleic acid_d17 (2 Everyvioleic	Cut. 10. 111 120-075	(Pac)-Indevimed (1-Methyl, DL trustenban) is an	
acid-d17) is the deuterium labeled		indoleamine 2,3-dioxygenase (IDO) inhibitor.	N
(Rac)-Idroxioleic acid. (Rac)-Idroxioleic acid (2-Hydroxyoleic acid) is a synthetic oleic acid	арарарар но		
(OA) derivative that binds to the plasma membrane	D D D D D D D D D D D D D D D D D D D		NH ₂
and alters lipid organization.		Durity 09.129/	Дон
Clinical Data: No Development Reported		Clinical Data: No Development Reported	0
Size: 1 mg, 5 mg		Size: 10 mM × 1 mL, 50 mg	

(S)-(+)-Rolipram		(S)-(–)-Perillyl alcohol	
((+)-Rolipram; (S)-Rolipram)	Cat. No.: HY-B0392		Cat. No.: HY-116514
(S)-(+)-Rolipram ((+)-Rolipram) is a cyclic AMP(cAMP)-specific phosphodiesterase 4 (PDE4) inhibitor, with an IC ₅₀ of 1100 nM. (S)-(+)-Rolipram can suppresse tumor necrosis factor-alpha (TNFα) production by human mononuclear cells. Purity: 99.89%		 (S)-(-)-Perillyl alcohol is a monoterpene found in lavender, inhibits farnesylation of Ras, upregulates the mannose-6-phosphate receptor and induces apoptosis. Anti-cancer activity. Purity: ≥98.0% 	С
Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg		Clinical Data: No Development Reported Size: 10 mM × 1 mL, 100 mg	
(S)-10-Hydroxycamptothecin (10-HCPT; 10-Hydroxycamptothecin)	Cat. No.: HY-N0095	(S)-10-Hydroxycamptothecin-d5 (10-HCPT-d5; 10-Hydroxycamptothecin-d5)	Cat. No.: HY-N0095S
(S)-10-Hydroxycamptothecin (10-HCPT;10-Hydroxycamptothecin) is a DNA topoisomerase I inhibitor of isolated from the Chinese plant Camptotheca accuminata.	HO-CJUN HO LO	(S)-10-Hydroxycamptothecin-d5 (10-HCPT-d5) is the deuterium labeled (S)-10-Hydroxycamptothecin. (S)-10-Hydroxycamptothecin (10-HCPT) is a DNA topoisomerase I inhibitor.	HO-GTN BHOTO
Purity: 99.38% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 50 mg, 100 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 10 mg	
(S)-Crizotinib	Cat. No.: HY-100549	(S)-Thalidomide ((S)-(-)-Thalidomide)	Cat. No. : HY-14658A
(S)-Crizotinib is a potent and selective MTH1 (mutT homologue) inhibitor with an IC _{so} of 330 nM.		(S)-Thalidomide ((S)-(-)-Thalidomide) is the S-enantiomer of Thalidomide. (S)-Thalidomide has immunomodulatory, anti-inflammatory, antiangiogenic and pro-apoptotic effects. (S)-Thalidomide induces teratogenic effects by binding to cereblop (CBBN)	
Purity: 99.61% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	0.0
(S)-Verapamil D7 hydrochloride	Cat No - HV 12522645	(S)-Verapamil hydrochloride	Cat No . HV 1252264
(S)-Verapamil D7 hydrochloride ((S)-(-)-Verapamil D7 hydrochloride) is a deuterium labeled (S)-Verapamil hydrochloride. (S)-Verapamil hydrochloride (S(-)-Verapamil hydrochloride) inhibits leukotriene C4 (LTC4) and calcein transport by MRP1.		(S)-Verapamil hydrochloride (S(-)-Verapamil hydrochloride) inhibits leukotriene C4 (LTC4) and calcein transport by MRP1. (S)-Verapamil hydrochloride leads to the death of potentially resistant tumor cells.	
Purity: >98% Clinical Data: No Development Reported Size: 1 mg		Purity: 99.39% Clinical Data: Launched Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
(S)-Verapamil-d6 hydrochloride ((S)-(-)-Verapamil-d6 hydrochloride)	Cat. No.: HY-135336AS1	(Z)-Guggulsterone	Cat. No. : HY-110066
(S)-Verapamil-d6 ((S)-(-)-Verapamil-d6) hydrochloride is the deuterium labeled (S)-Verapamil hydrochloride. (S)-Verapamil hydrochloride (S(-)-Verapamil hydrochloride) inhibits leukotriene C4 (LTC4) and calcein transport by MRP1.		Z-guggulsterone, a constituent of Indian Ayurvedic medicinal plant Commiphora mukul, inhibits the growth of human prostate cancer cells by causing apoptosis . Z-guggulsterone inhibits angiogenesis by suppressing the VEGF-VEGF-R2-Akt signaling axis.	
Purity. >98% Clinical Data: No Development Reported Size: 1 mg, 10 mg		Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	00 mg

(Z)-SU5614		(±)-Evodiamine	
(Z)-SU5614 is a potent FLT3 inhibitor and selectively induces growth arrest, apoptosis, and cell cycle arrest in Ba/F3 and AML cell lines expressing a constitutively activated FLT3.	Cat. No.: HY-18952A	(±)-Evodiamine, a quinazolinocarboline alkaloid, is a Top1 inhibitor. Evodiamine exhibits anti-inflammatory, antiobesity, and antitumor effects. (±)-Evodiamine inhibits the proliferation of a wide variety of tumor cells by inducing their apoptosis	Cat. No.: HY-N0114A
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	/	Purity:99.97%Clinical Data:No Development ReportedSize:250 mg, 500 mg, 1 g	1355
(±)-Norcantharidin ((±)-NCTD)	Cat. No. : HY-N0291	1-(4-Chloro-3-(trifluoromethyl)phenyl)-3-(4-(4-	cyanophenoxy) Cat. No.: HY-136658
(±)-Norcantharidin ((±)-NCTD) is a compound possessing anti-angiogenetic activity with potential use in anti-cancertherapy.		STAT3-IN-7 is a Sorafenib analogue and potently inhibits the phosphorylation of STAT3 . STAT3-IN-7 induces cell apoptosis through SHP-1 dependent STAT3 inactivation. STAT3-IN-7 does not inhibit kinase activity and has anticancer effects.	N°C°C [#] L [#] C [°] _F
Purity:>98%Clinical Data:No Development ReportedSize:5 mg	H.	Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	
1-Alaninechlamydocin	Cat. No.: HY-P2698	12-HETE	Cat. No.: HY-113439
 1-Alaninechlamydocin, a cyclic tetrapeptide, is a potent HDAC inhibitor (IC₅₀=6.4 nM). 1-Alaninechlamydocin induces G2/M cell cycle arrest and apoptosis in MIA PaCa-2 cells. Purity: >98% Clinical Data: No Development Reported 		12-HETE, a major metabolic product of arachidonic acid using 12-LOX catalysis, inhibits cell apoptosis in a dose-dependent manner. 12-HETE promotes the activation and nuclear translocation of NF-kB through the integrin-linked kinase (ILK) pathway. Purity: >98% Clinical Data: No Development Reported	on States
Size: 1 mg, 5 mg		Size: 100 µg	
12-HETE-d8	Cat. No.: HY-113439S	13-Methyltetradecanoic acid (13-MTD; 13-Methylmyristic acid)	Cat. No.: HY-131503
12-HETE-d8 is the deuterium labeled 12-HETE. 12-HETE, a major metabolic product of arachidonic acid using 12-LOX catalysis, inhibits cell apoptosis in a dose-dependent manner.		13-Methyltetradecanoic acid (13-MTD), a saturated branched-chain fatty acid with potent anticancer effects. 13-Methyltetradecanoic acid induces apoptosis in many types of human cancer cells.	Land Carl
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	7	Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg	
14215	Cat. No.: HY-139865	15-Acetoxyscirpenol	Cat. No.: HY-N6681
14215 is a potent ML-IAP Lys-covalent inhibitor with an $\rm IC_{50}$ value of 11 nM.		15-acetoxyscirpenol, one of acetoxyscirpenol moiety mycotoxins (ASMs), strongly induces apoptosis and inhibits Jurkat T cell growth in a dose-dependent manner by activating other caspases independent of caspase-3.	Н.О.Н. ОН ОГООН
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	. 4.	Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	1970) 1970

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3-Dehydrotrametenolic acid		3-Hydroxykynurenine	
	Cat. No.: HY-N2177	(3-Hydroxy-DL-kynurenine)	Cat. No.: HY-113294
3-Dehydrotrametenolic acid, isolated from the sclerotium of Poria cocos, is a lactate dehydrogenase (LDH) inhibitor. 3-Dehydrotrametenolic acid promotes adipocyte differentiation in vitro and acts as an insulin sensitizer in vivo.	ностран	 3-Hydroxykynurenine, a metabolite of tryptophan, is a potential endogenous neurotoxin whose increased levels have been described in several neurodegenerative disorders. 3-Hydroxykynurenine induces neuronal apoptosis. 	HO NH2 O NH2
Purity:99.86%Clinical Data:No Development ReportedSize:5 mg, 10 mg	In The second seco	Purity: ≥98.0% Clinical Data: No Development Reported Size: 5 mg, 10 mg	
3-Hydroxykynurenine-d3 (3-Hydroxy-DL-kynurenine-d3)	Cat. No.: HY-113294S1	3-Hydroxyterphenyllin (NSC 299113)	Cat. No .: HY-N10268
3-Hydroxykynurenine-d3 (3-Hydroxy-DL-kynurenine-d3) is the deuterium labeled 3-Hydroxykynurenine.		3-Hydroxyterphenyllin is a metabolite of Aspergillus candidus.3-Hydroxyterphenyllin suppresses proliferation and causes cytotoxicity against A2780/CP70 and OVCAR-3 cells. 3-Hydroxyterphenyllin induces S phase arrest and apoptosis .	HO, CH, OH HO, CH, OH, O, CH, OH, OH, OH, OH, OH, OH, OH, OH, OH, O
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
2 Nitropropapois asid		2.0 Mothylgallic acid	
(β-Nitropropionic acid; Bovinocidin)	Cat. No.: HY-W012875	(3,4-Dihydroxy-5-methoxybenzoic acid)	Cat. No.: HY-N2009
3-Nitropropanoic acid (β -Nitropropionic acid) is an irreversible inhibitor of succinate dehydrogenase . 3-Nitropropanoic acid exhibits potent antimycobacterial activity with a MIC value of 3.3 μ M.	O −O ^{−N*} O	3-O-Methylgallic acid (3,4-Dihydroxy-5-methoxybenzoic acid) is an anthocyanin metabolite and has potent antioxidant capacity. 3-O-methylgallic acid inhibits Caco-2 cell proliferation with an IC_{50} value of 24.1 μ M.	но си
Purity:99.93%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 100 mg, 500 mg		Purity:97.76%Clinical Data:No Development ReportedSize:500 mg, 1 g	ОН
3-O-Acetyloleanolic acid		3BDO	
	Cat. No.: HY-N2618		Cat. No.: HY-U00434
3-O-Acetyloleanolic acid (3AOA), an oleanolic acid derivative isolated from the seeds of Vigna sinensis K., induces in cancer and also exhibits anti-angiogenesis activity.	Loc H I Dot	3BDO is a new mTOR activator which can also inhibit autophagy .	
Purity:> 98%Clinical Data:No Development ReportedSize:5 mg, 10 mg		Purity:99.91%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 25 mg	
4'-Bromo-resveratrol (4'BR)	Cat. No.: HY-124113	4-Bromo A23187	Cat. No.: HY-N6694
4'-Bromo-resveratrol is a potent and dual inhibitor Sirtuin-1 and Sirtuin-3 . 4'-Bromo-resveratrol inhibits melanoma cell growth through mitochondrial metabolic reprogramming.	Br OH	4-Bromo A23187 is a halogenated analog of the highly selective calcium ionophore A-23187. 4-Bromo A23187a calcium modulator, induces apoptosis in different cells, including HL-60 cells.	
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	Un	Purity:≥99.0%Clinical Data:No Development ReportedSize:1 mg	

4-Hydroperoxy cyclophosphamide	Cat. No.: HY-117433	4-Hydroperoxy Cyclophosphamide-d4	Cat. No.: HY-117433S
4-Hydroperoxy cyclophosphamide is the active metabolite form of the prodrug Cyclophosphamide.	CI ∕ NH O ^{SP} I CI ∕ N CI	 4-Hydroperoxy Cyclophosphamide-d4 is the deuterium labeled 4-Hydroperoxy cyclophosphamide. 4-Hydroperoxy cyclophosphamide is the active metabolite form of the prodrug Cyclophosphamide. 	
Clinical Data: No Development Reported Size: 5 mg, 10 mg		Clinical Data: Size: 1 mg, 5 mg	
4-Hydroxybenzyl alcohol	Cat. No. : HY-Y0892	4-Methyldaphnetin	Cat. No.: HY-N4286
4-Hydroxybenzyl alcohol is a phenolic compound widely distributed in various kinds of plants. Anti-inflammatory, anti-oxidant, anti-nociceptive activity. Neuroprotective effect. Inhibitor of tumor angiogenesis and growth.	НООН	 4-Methyldaphnetin is a precursor in the synthesis of derivatives of 4-methyl coumarin. 4-Methyldaphnetin has potent, selective anti-proliferative and apoptosis-inducing effects on several cancer cell lines. 	HO OH OF O
Purity:99.34%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 100 mg		Purity:99.43%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 20 mg	
4-MMPB	Cat. No. : HY-118480	4-tert-Octylphenol	Cat. No.: HY-B1941
4-MMPB is a selective inhibitor of 15-lipoxygenase , with an IC_{50} of 18 μ M. 4-MMPB has IC_{50} s of 19.5 μ M and 19.1 μ M for soybean 15-lipoxygenase (SLO) and human 15-lipoxygenase-1 (15-LOX-1), respectively. 4-MMPB has potential for the research of prostate cancer.	S N N N N N	4-tert-Octylphenol, a endocrine-disrupting chemical, is an estrogenic drug. 4-tert-Octylphenol induces apoptosis in neuronal progenitor cells in offspring mouse brain.	<u>а</u> рун
Purity: 99.69% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	00 mg	Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	
4-Vinylphenol	Cat. No.: HY-W005288	4EGI-1	Cat. No.: HY-19831
4-Vinylphenol is found in the medicinal herb Hedyotis diffusa Willd, wild rice and is also the metabolite of p-coumaric and ferulic acid by lactic acid bacteria in wine.	HO	4EGI-1 is an inhibitor of $eIF4E/eIF4G$ interaction, with a K_a of 25 μM against eIF4E binding.	о, , , , , , , , , , , , , , , , , , ,
Purity: >98% Clinical Data: No Development Reported Size: 100 mg (832.2 mM * 1 mL in Propylene glycol)),	Purity: 98.83% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	00 mg
5'-Methylthioadenosine (5'-(Methylthio)-5'-deoxyad 5'-Deoxy-5'-(methylthio)adenosine;)	enosine; Cat. No.: HY-16938	5,8-Epidioxyergosta-6,9(11),22-trien-3-ol (9,11-Dehydroergosterol peroxide; 9(11)-DHEP)	Cat. No. : HY-N7175
5'-Methylthioadenosine (5'-(Methylthio)-5'-deoxyadenosine) is a nucleoside generated from S-adenosylmethionine (SAM) during polyamine synthesis.		5,8-Epidioxyergosta-6,9(11),22-trien-3-ol (9,11-Dehydroergosterol peroxide), an important steroid from medicinal mushroom, exerts antitumor activity in several tumor types.	HO LOS H
Purity:99.67%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 25 mg	HO	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	- 12

5,7,4'-Trimethoxyflavone Cat. No.: HY-N6818	5-(N,N-Hexamethylene)-amiloride (Hexamethylene amiloride; HMA) Cat. No.: HY-128067
5,7,4'-Trimethoxyflavone is isolated from Kaempferia parviflora (KP) that is a famous medicinal plant from Thailand.	5-(N,N-Hexamethylene)-amiloride (Hexamethylene amiloride) derives from an amiloride and is a potent Na ⁺ /H ⁺ exchanger inhibitor, which decreases the intracellular pH (pH _i) and induces apoptosis in leukemic cells.
Purity:99.78%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 20 mg	Purity: 98.42% Clinical Data: Phase 3 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg
5-Aminolevulinic acid hydrochloride (5-ALA hydrochloride; δ-Aminolevulinic acid hydrochloride;) Cat. No.: HY-N0305	5-Aminolevulinic acid-15N hydrochloride (5-ALA-15N hydrochloride;) Cat. No.: HY-N0305S
5-Aminolevulinic acid hydrochloride (5-ALA hydrochloride) is an intermediate in heme biosynthesis in the body and the universal precursor of tetrapyrroles. H ₂ N H-Cl	5-Aminolevulinic acid-15N (5-ALA-15N)hydrochloride is the 15N-labeled 5-Aminolevulinicacid (hydrochloride). 5-Aminolevulinic acidhydrochloride (5-ALA hydrochloride) is anintermediate in heme biosynthesis in the body andthe universal precursor of tetrapyrroles.
Purity: >97.0% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 1 g, 5 g, 10 g	Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg
E Aminologulinis acid d2 hydrochlarida (r. 414-42	E Eluprovracil
hydrochloride;) Cat. No.: HY-N030551	(5-FU) Cat. No.: HY-90006
5-Aminolevulinic acid-d2 (hydrochloride) is deuterium labeled 5-Aminolevulinic acid (hydrochloride).	5-Fluorouracil (5-FU) is an analogue of uracil and a potent antitumor agent. 5-Fluorouracil affects pyrimidine synthesis by inhibiting thymidylate synthetase thus depleting intracellular dTTP pools. 5-Fluorouracil induces apoptosis and can be used as a chemical sensitiver
HCI Purity: >98% Clinical Data: Size: 1 mg, 5 mg	Purity: 99.86% Clinical Data: Launched Size: 10 mM × 1 mL, 200 mg, 1 g, 5 g
5-Fluorouracii-15N2 Cat. No.: HY-90006S2	5-Fluorouracil-d1 (5-FU-d1) Cat. No.: HY-90006S
5-Fluorouracil-15N2 is the 15N-labeled 5-Fluorouracil. 5-Fluorouracil (5-FU) is an analogue of uracil and a potent antitumor agent. 5-Fluorouracil affects pyrimidine synthesis by inhibiting thymidylate synthetase thus depleting intracellular dTTP pools. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	5-Fluorouracil-d1 (5-FU-d1) is the deuterium labeled 5-Fluorouracil. 5-Fluorouracil (5-FU) is an analogue of uracil and a potent antitumor agent. 5-Fluorouracil affects pyrimidine synthesis by inhibiting thymidylate synthetase thus depleting intracellular dTTP pools. Purity: >98% Clinical Data: No Development Reported Size: 5 mg, 50 mg
5β-Dihydrocortisol Cat. No.: HY-N3995	5β-Dihydrocortisol-d6 Cat. No.: HY-N3995S
5 β -Dihydrocortisol, a metabolite of Cortisol, is a potential mineralocorticoid. 5 β -Dihydrocortisol can potentiate glucocorticoid activity in raising the intraocular pressure. 5 β -Dihydrocortisol causes breast cancer cell apoptosis.	5β-Dihydrocortisol-d6 is the deuterium labeled 5β-Dihydrocortisol, 5β-Dihydrocortisol, a metabolite of Cortisol, is a potential mineralocorticoid. 5β-Dihydrocortisol can potentiate glucocorticoid activity in raising the intraocular pressure.
Purity: 98.05% Clinical Data: No Development Reported Size: 10 mM × 1 mL 5 mg, 10 mg, 25 mg	Purity: >98% Clinical Data: No Development Reported Size: 1 ma. 5 ma



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9-ING-41		9-Methoxycamptothecin	
	Cat. No.: HY-113914		Cat. No.: HY-N6011
9-ING-41 is a maleimide-based ATP-competitive and selective glycogen synthase kinase-3 β (GSK-3 β) inhibitor with an IC _{so} of 0.71 μ M. 9-ING-41 significantly leads to cell cycle arrest, autophagy and apoptosis in cancer cells.	HN HO N HO	 9-Methoxycamptothecin (MCPT), isolated from Nothapodytes foetida, has antitumor activities through topoisomerase inhibition. 9-Methoxycamptothecin (MCPT) induces strong G2/M arrest and apoptosis in cancer. 	SHOT N N N N
Clinical Data: Phase 2	r	Clinical Data: No Development Reported	
Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	100 mg	Size: 5 mg, 10 mg, 20 mg	
	5	J. J. J	
A 1010477		A 102621	
A-1210477	Cat No. UV 12469	A-192021	Cat No. UV 12020E
	Cat. No.: HY-12468		Cat. NO.: HY-120295
A-1210477 is a potent and selective inhibitor of $MCL-1$ with a K_i of 0.45 nM. A-1210477 specifically binds MCL-1 and promotes apoptosis of cancer cells in an MCL-1-dependent manner.	All and a second s	A-192621 is a potent, nonpeptide, orally active and selective endothelin B (ET_e) receptor antagonist with an IC _{so} of 4.5 nM and a K ₁ of 8.8 nM. The selectivity of A-192621 is 636-fold higher than ET _A (IC _{so} of 4280 nM and K ₁ of 5600 nM). A-192621 promotes apoptosis in PASMCs.	- HN-C H CHS
Purity: 98.89%	~~~	Purity: 99.85%	
Clinical Data: No Development Reported		Clinical Data: No Development Reported	
Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg		Size: 5 mg	
AAPK-25		Abacavir	
	Cat. No.: HY-126249		Cat. No.: HY-17423
AAPK-25 is a potent and selective Aurora/PLK dual inhibitor with anti-tumor activity, which can cause mitotic delay and arrest cells in a prometaphase, reflecting by the biomarker histone H3 ^{Ser10} phosphorylation and followed by a surge	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Abacavir is a potent nucleoside analog reverse-transcriptase inhibitor (NRTI).	
in apoptosis. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Purity: 99.70% Clinical Data: Launched Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg	,, N=/
Abacavir sulfate		Abacavir-d4	
(Abacavir Hemisulfate; ABC sulfate)	Cat. No.: HY-17423A		Cat. No.: HY-17423S
Abacavir sulfate (ABC) is a powerful nucleoside analog reverse transcriptase inhibitor (NRTI) used to treat HIV and AIDS.		Abacavir-d4 is the deuterium labeled Abacavir. Abacavir is a potent nucleoside analog reverse-transcriptase inhibitor (NRTI) .	H2N N N N N N N N
	18 0-8-04		D NH
Purity: 99.81%	OH OH	Purity: >98%	X
Clinical Data: Launched		Clinical Data: No Development Reported	00
Size: 10 mM × 1 mL, 10 mg, 50 mg		Size: 1 mg, 5 mg	
ABL-L		ABT 737-d8	
	Cat. No.: HY-142913		Cat. No.: HY-50907S
API Linduce anotheric of human langer and		APT 727 dQ is the douterium labeled APT 727	
ABL-L induces apoptosis of numan laryngocarcinoma cells through p53-dependent pathway.	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	ABT 737-08 Is the deuterium labeled ABT-737. ABT-737, a BH3 mimetic, is a potent Bcl-2 , Bcl-x _L and Bcl-w inhibitor with EC _{so} s of 30.3 nM, 78.7 nM, and 197.8 nM, respectively.	
Purity: >98%	0	Purity: >98%	1~~~Q
Clinical Data: No Development Reported		Clinical Data: No Development Reported	
Size: 1 mg, 5 mg		Size: 1 mg, 10 mg	

ABT-100		ABT-737	
	Cat. No.: HY-119257		Cat. No.: HY-50907
ABT-100 is a potent, highly selective and orally active farnesyltransferase inhibitor.	N O OHI	ABT-737, a BH3 mimetic, is a potent Bcl-2 , Bcl-x _L and Bcl-w inhibitor with EC ₅₀ s of 30.3 nM, 78.7 nM, and 197.8 nM, respectively. ABT-737 induces the disruption of the BCL-2/BAX complex and BAK-dependent but BIM-independent activation of the intrinsic apoptotic pathway.	ju nitori
Purity:98.18%Clinical Data:No Development ReportedSize:5 mg	F∱F	Purity: 99.72% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	200 mg
Acacetin		ACBI1	
(5,7-Dihydroxy-4'-methoxyflavone)	Cat. No.: HY-N0451		Cat. No.: HY-128359
Acacetin (5,7-Dihydroxy-4'-methoxyflavone) is an orally active flavonoid derived from Tephroseris kirilowii (Turcz.) Holub. Acacetin docks in the ATP binding pocket of PI3Ky. Acacetin causes cell cycle arrest and induces apoptosis and autophagy in cancer cells.		ACBI1 is a potent PROTAC degrader of BAF ATPase subunits SMARCA2 and SMARCA4 , also degrades the polybromo-associated BAF (PBAF) complex member PBRM1 , with DC ₅₀ S of 6 nM, 11 nM and 32 nM for SMARCA2, SMARCA4 and PBRM1 in MV-4-11 cells, respectively.	
Clinical Data: No Development Reported		Clinical Data: No Development Reported	
Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg		Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
Acetylcysteine		Acetylcysteine-15N	
(N-Acetylcysteine; N-Acetyl-L-cysteine; NAC)	Cat. No.: HY-B0215	(N-Acetylcysteine-15N; N-Acetyl-L-cysteine-15N; NAC-15	N) Cat. No.: HY-B0215S1
Acetylcysteine (N-Acetylcysteine) is a mucolytic agent which reduces the thickness of the mucus. Acetylcysteine is a ROS inhibitor.	о М Сон	Acetylcysteine-15N (N-Acetylcysteine-15N) is the 15N-labeled Acetylcysteine. Acetylcysteine (N-Acetylcysteine) is a mucolytic agent which reduces the thickness of the mucus. Acetylcysteine is a ROS inhibitor.	
Purity:≥98.0%Clinical Data:LaunchedSize:500 mg, 5 g, 10 g	" 0	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	 0
Asstukavetsing d2		Acitrotia	
(N-Acetylcysteine-d3: N-Acetyl-L-cysteine-d3: NAC-d3)		(Ro 10-1670)	
	Cat. No 111-002155		Cat. No.: 111-00107
Acetylcysteine-d3 (N-Acetylcysteine-d3) is the deuterium labeled Acetylcysteine. Acetylcysteine (N-Acetylcysteine) is a mucolytic agent which reduces the thickness of the mucus. Acetylcysteine is a ROS inhibitor.		Acitretin (Ro 10-1670) is a second-generation, systemic retinoid that has been used in the treatment of psoriasis. Acitretin also can be used for the research of Alzheimer's disease.	, fiched in
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	SH	Purity: 99.79% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 200 mg, 500 mg	
Asitustia d2		ACT201	
Actireun-05 (Ro 10-1670-d3)	Cat. No.: HY-B0107S	ACIUUI	Cat. No.: HY-128861A
Acitretin-d3 (Ro 10-1670-d3) is the deuterium labeled Acitretin. Acitretin (Ro 10-1670) is a second-generation, systemic retinoid that has been used in the treatment of psoriasis. Acitretin also can be used for the research of Alzheimer's disease.	of other stands on	ACT001 is an orally active PAI-1 inhibitor by inhibiting the phosphorylation of PI3K and AKT . ACT001 inhibits the phosphorylation of STAT3 and PD-L1 expression by directly binding to STAT3 .	ног и страние но стори
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Purity: 99.62% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg	

Actein	Cot No . UV N6972	Actinonin	Cat. No . UV 112052
Actein is a triterpene glycoside isolated from the rhizomes of Cimicifuga foetida. Actein suppresses cell proliferation, induces autophagy and apoptosis through promoting ROS/JNK activation, and blunting AKT pathway in human bladder cancer. Actein has little toxicity in vivo. Purity : 98.58% Clinical Data: No Development Reported Size: 5 mg		((-)-Actinonin) Actinonin ((-)-Actinonin) is a naturally occurring antibacterial agent produced by Actinomyces. Actinonin inhibits aminopeptidase M, aminopeptidase N and leucine aminopeptidase. Purity: 99.30% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg,	на р р р р р р р р р р р р р р р р р р р
Acyclovir (Aciclovir; Acycloguanosine)	Cat. No.: HY-17422	Acyclovir-d4 (Aciclovir-d4; Acycloguanosine-d4)	Cat. No.: HY-17422S1
Acyclovir (Aciclovir) is a guanosine analogue and an orally active antiviral agent. Acyclovir inhibits HSV-1 (IC ₅₀ of 0.85 μ M), HSV-2 (IC ₅₀ of 0.86 μ M) and varicella-zoster virus.		Acyclovir-d4 (Aciclovir-d4) is the deuterium labeled Acyclovir. Acyclovir (Aciclovir) is a guanosine analogue and an orally active antiviral agent. Acyclovir inhibits HSV-1 (IC ₅₀ of 0.85 μ M), HSV-2 (IC ₅₀ of 0.86 μ M) and varicella-zoster virus.	
Purity: 99.34% Clinical Data: Launched Size: 10 mM × 1 mL 50 mg 100 mg 500 mg	0H	Purity: >98% Clinical Data: No Development Reported Size: 1 mg 5 mg	- 01
		5126. 1 mg, 5 mg	
Acyclovir-d4 L-Leucinate		Adapalene	
	Cat. No.: HY-17422S	(CD271)	Cat. No.: HY-B0091
Acyclovir-d4 L-Leucinate is the deuterium labeled Acyclovir. Acyclovir (Aciclovir) is a guanosine analogue and an orally active antiviral agent. Acyclovir inhibits HSV-1 (IC ₅₀ of 0.85 μ M), HSV-2 (IC ₅₀ of 0.86 μ M) and varicella-zoster virus.	Ante o o the second	Adapalene (CD271), a third-generation synthetic retinoid, is widely used for the research of acne. Adapalene is a potent RAR agonist, with $AC_{so}s$ of 2.3 nM, 9.3 nM, and 22 nM for RAR β , RAR γ , RAR α , respectively.	HOY
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 10 mg		Purity: ≥98.0% Clinical Data: Launched Size: 10 mM × 1 mL, 50 mg, 100 mg, 500 mg	
Adapalene sodium salt		Adapalene-d3	
(CD 271 sodium salt)	Cat. No.: HY-B0091A		Cat. No.: HY-B0091S
Adapalene (CD271) sodium salt, a third-generation synthetic retinoid, is widely used for the research of acne. Adapalene sodium salt is a potent RAR agonist, with AC_{50} s of 2.3 nM, 9.3 nM, and 22 nM for RAR β , RAR γ , RAR α , respectively. Purity: >98% Clinical Data: Launched Size: 1 mg, 5 mg	App COS ^{Lo} No.	Adapalene-d3 is the deuterium labeled Adapalene. Adapalene (CD271), a third-generation synthetic retinoid, is widely used for the research of acne. Adapalene is a potent RAR agonist, with AC ₅₀ S of 2.3 nM, 9.3 nM, and 22 nM for RAR β , RAR γ , RAR α , respectively. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 10 mg	С С С С С С С С С С С С С С С С С С С
Adapalene-d6 Methyl Ester	Cat. No.: HY-B0091S1	Adaphostin (NSC 680410)	Cat. No.: HY-103275
Adapalene-d6 Methyl Ester is the deuterium labeled Adapalene. Adapalene (CD271), a third-generation synthetic retinoid, is widely used for the research of acne. Adapalene is a potent RAR agonist, with AC _{so} s of 2.3 nM, 9.3 nM, and 22 nM for RAR β , RAR γ , RAR α , respectively.	OF Street	Adaphostin (NSC 680410), the adamantyl ester of AG957, is a potent p210 ^{bcr/abl} inhibitor (IC_{50} =14 μ M). Adaphostin induces apoptosis in T-lymphoblastic human leukemia cell lines (IC_{50} ranging from 17 to 216 nM).	HOLICH
Purity: > 98% Clinical Data: No Development Reported Size: 10 mg, 100 mg	×₽ ₽ ₽	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	



AG-024322		AG-825	
	Cat. No.: HY-15491	(Tyrphostin AG-825)	Cat. No.: HY-15844
AG-024322 is a potent ATP-competitive pan-CDK inhibitor against cell cycle kinases CDK1, CDK2, and CDK4 with K _i values in the 1-3 nM range. AG-024322 displays broad-spectrum anti-tumor activity and clear target modulation in vivo. AG-024322 induces cell apoptosis . Purity: 98.69% Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg, 50 mg		AG-825 (Tyrphostin AG-825) is a selective and ATP-competitive ErbB2 inhibitor which suppresses tyrosine phosphorylation, with an IC_{50} of 0.35 μ M. AG-825 displays anti-cancer activity. AG825 significantly accelerates apoptosis of human neutrophils.Purity:98.07%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	(1 + 1) + 1 + 1 + 1 + 1 + 1 + 1 + 1 + 1 +
AG1024	C + N - 11/ 10050	AG6033	C + N - UV 142425
(Tyrphostin AG 1024)	Cat. No.: HY-10253		Cat. No.: HY-143435
$\begin{array}{llllllllllllllllllllllllllllllllllll$	HO	AG6033 is a potential novel CRBN modulator. AG6033 suppresses various tumor cells by modulating the interactions between CRBN and various antitumor target proteins. AG6033 can cause GSPT1 and IKZF1 degradation. AG6033 induces CRBN -dependent cytotoxic effect. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	CT-CC N-NH of CD NNNH of CD NNNH of CD NNNH O
AGK2		AGN194204	
	Cat. No.: HY-100578	(IRX4204; NRX194204; VTP 194204)	Cat. No.: HY-13717
AGK2 is a selective SIRT2 inhibitor with an IC ₅₀ of 3.5 μ M. AGK2 inhibits SIRT1 and SIRT3 with IC ₅₀ s of 30 and 91 μ M, respectively.		AGN194204 (IRX4204) is an orally active and selective RXR agonist with K_d values 0.4 nM, 3.6 nM and 3.8 nM and EC _{sp} s of 0.2 nM, 0.8 nM and 0.08 nM for RXR α , RXR β and RXR γ , respectively. AGN194204 is inactive against RAR.	And and an
Purity: 99.62% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	0 mg	Purity: ≥99.0% Clinical Data: Phase 2 Size: 1 mg, 5 mg	
Ajoene	C + N + 10(100704	AK-778-XXMU	C . N. 19/ 144707
	Cat. No.: HY-106784		Cat. No.: HY-144707
Ajoene, a garlic-derived compound, is an antithrombotic and antifungal agent. Ajoene inhibits proliferation and induces apoptosis of human leukaemia CD34-negative cells including HL-60, U937, HEL and OCIM-I. Anticancer activities. Purity: >98%	0 \$~~\$~~\$. \$~~\$	AK-778-XXMU is a potent inhibitor of DNA Binding 2 (ID2) antagonist with a K _p of 129 nM. AK-778-XXMU can inhibit cell migration and invasion of glioma cell lines, induce apoptosis , and more importantly, slow down the tumor growth.	CI HO HO N
Clinical Data: No Development Reported		Clinical Data: No Development Reported	
Size: 5 mg, 10 mg, 25 mg		Size: 1 mg, 5 mg	
AKT inhibitor VIII		AKT-IN-3	
(AKTi-1/2)	Cat. No.: HY-10355		Cat. No.: HY-126257
AKT inhibitor VIII (AKTi-1/2) is a cell-permeable quinoxaline compound that has been shown to potently, selectively, allosterically, and reversibly inhibit Akt1, Akt2, and Akt3 activity with IC ₅₀ s of 58 nM, 210 nM, and 2119 nM, respectively.	Frank and	AKT-IN-3 (compound E22) is a potent, orally active low hERG blocking Akt inhibitor, with 1.4 nM, 1.2 nM and 1.7 nM for Akt1, Akt2 and Akt3, respectively.	
Purity: 98.93% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 50 mg, 100 mg		Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg	
	www.MedCh	lemExpress.com	1



alpha-Bisabolol		alpha-Hederin	
alpha-Bisabolol is a nontoxic sesquiterpene alcohol present in natural essential oil, with anticancer activity.	Сат. No.: HY-121222	$(\alpha$ -Hederin) alpha-Hederin (α -Hederin), a monodesmosidic triterpenoid saponin, exhibits promising antitumor potential against a variety of human cancer cell lines.	Cat. No.: HY-NU255
Purity: ≥80.0% Clinical Data: No Development Reported Size: 500 mg, 1 g		Purity: ≥98.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg	0 ⁻² CH
alpha-Mangostin (α-Mangostin)	Cat. No.: HY-N0328	Alrizomadlin (APG-115; AA-115)	Cat. No.: HY-101518
alpha-Mangostin (α -Mangostin) is a dietary xanthone with broad biological activities, such as antioxidant, anti-allergic, antiviral, antibacterial, anti-inflammatory and anticancer effects. It is an inhibitor of mutant IDH1 (IDH1-R132H) with a K ₁ of 2.85 μ M. Purity: 99.64% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 25 mg, 50 mg, 100 mg		Alrizomadlin (APG-115) is an orally active MDM2 protein inhibitor binding to MDM2 protein with IC ₅₀ and K _i values of 3.8 nM and 1 nM, respectively. Alrizomadlin blocks the interaction of MDM2 and p53 and induces cell-cycle arrest and apoptosis in a p53-dependent manner. Purity: 98.16% Clinical Data: Phase 2 Size: 1 mg, 5 mg, 10 mg	
Alsterpaulione		Alteminostat	
(9-Nitropaulione; NSC 705701) Alsterpaulione (9-Nitropaulione) is a potent CDK inhibitor, with IC ₅₀ s of 35 nM, 15 nM, 200 nM and 40 nM for CDK1/cyclin B, CDK2/cyclin A, CDK2/cyclin E and CDK5/p35, respectively. Purity: 98.38% Clinical Data: No Development Reported Size: 5 mg, 10 mg	-0-N [*] 0	Alteminostat (CKD-581) is a potent HDAC inhibitor. Alteminostat inhibits the class I-II HDAC family via histone H3 and tubulin acetylation. Alteminostat can be used for lymphoma and multiple myeloma research. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	Сат. No.: HY-109109
Alvespimycin hydrochloride (17-DMAG hydrochloride; KOS-1022; BMS 826476)	Cat. No .: HY-12024	Amarogentin	Cat. No.: HY-N2447
Alvespimycin hydrochloride (17-DMAG hydrochloride; KOS-1022; BMS 826476) is a potent inhibitor of Hsp90, binding to Hsp90 with EC_{50} of 62±29 nM.		Amarogentin is a secoiridoid glycoside that is mainly extracted from Swertia and Gentiana roots. Amarogentin exhibits many biological effects, including anti-oxidative, anti-tumour, and anti-diabetic activities.	но орности орно орно орно орности орно орно орно орно орно орно орно орн
Purity: 98.68% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 10 mg, 25 mg, 100 mg, 200 mg	на	Purity: 98.96% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	о с _{он}
Amentoflavone (Didemethyl-ginkgetin)	Cat. No.: HY-N0662	Amiloride (MK-870)	Cat. No.: HY-B0285
Amentoflavone is a natural biflavone compound with many biological properties, including anti-inflammatory, antioxidative, and neuroprotective effects.	но с с с с с с с с с с с с с с с с с с с	Amiloride (MK-870) is an inhibitor of both epithelial sodium channel (ENaC) and urokinase-type plasminogen activator receptor (uTPA). Amiloride is a blocker of polycystin-2 (PC2; TRPP2) channel.	
Purity:98.88%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	он о	Purity:>98%Clinical Data:LaunchedSize:1 mg, 5 mg	

Amiloride hydrochloride		Amiloride hydrochloride dihydrate	
(MK-870 hydrochloride)	Cat. No.: HY-B0285A	(MK-870 hydrochloride dihydrate)	Cat. No.: HY-B0285B
Amiloride hydrochloride (MK-870 hydrochloride) is an inhibitor of both epithelial sodium channel (ENaC) and urokinase-type plasminogen activator receptor (uTPA). Amiloride hydrochloride is a blocker of polycystin-2 (PC2; TRPP2) channel. Purity: 99.65% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 500 mg	$\begin{array}{c} CI \\ H_2N \end{array} N \\ H_2N \end{array} H \\ CI \\ H_2 N \\ N \\ H_2 \\ HCI \\ HCI \\ \end{array} \begin{array}{c} NH \\ H_2 \\ HH_2 \\ HCI \\ HC$	Amiloride hydrochloride dihydrate (MK-870 hydrochloride dihydrate) is an inhibitor of both epithelial sodium channel (ENaC) and urokinase-type plasminogen activator receptor (uTPA). Amiloride hydrochloride dihydrate is a blocker of polycystin-2 (PC2; TRPP2) channel. Purity: 99.70% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg	$\begin{array}{c} CI \\ H_2N \\ H_2N \\ H_2N \\ H_2 \\ H_2$
Aminopurvalanol A	Cat. No.: HY-104013	Amsilarotene (TAC-101; Am 555S)	Cat. No.: HY-14653
Aminopurvalanol A is a potent, selective, and cell permeable inhibitor of Cyclins/Cdk complexes. Aminopurvalanol A preferentially targets the G2/M-phase transition inhibiting cancer cell differentiation. Purity: 98.00%		Amsilarotene (TAC-101; Am 555S), an orally active synthetic retinoid, has selective affinity for retinoic acid receptor α (RAR-α) binding with K_i of 2.4, 400 nM for RAR-α and RAR-β. Purity: 99.70%	Sa Charles Cont
Clinical Data: No Development Reported		Clinical Data: No Development Reported	
Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg		Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
Amuvatinib		AMXT-1501 tetrahydrochloride	
(MP470; HPK 56)	Cat. No.: HY-10206		Cat. No.: HY-124617A
Amuvatinib (MP470) is an orally bioavailable multi-targeted tyrosine kinase inhibitor with potent activity against mutant c-Kit , PDGFRα , Flt3 , c-Met and c-Ret .	S NH CS	AMXT-1501 tetrahydrochloride is an orally active polyamine transport inhibitor. AMXT1501 blocks tumor growth in immunocompetent mice but not in athymic nude mice lacking T cells. Combination of DFMO and AMXT1501 induces caspase3 mediated apoptosis in NB cell lines.	
Purity: 98.07% Clinical Data: Phase 2	C∕ N [⊥]	Purity: ≥98.0% Clinical Data: Phase 1	
Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Size: 5 mg, 10 mg, 25 mg, 50 mg	
Angelicin		Angiotensin II human	
(Isopsoralen)	Cat. No.: HY-N0763	(Angiotensin II; Ang II; DRVYIHPF)	Cat. No.: HY-13948
Angelicin, a furocoumarin naturally occurring tricyclic aromatic compound, structurally related to psoralens, is reported to have anti-cancer, antiviral, anti-inflammatory activity.	0	Angiotensin II (Angiotensin II) is a vasoconstrictor and a major bioactive peptide of the renin/angiotensin system.	to the second
Purity:99.86%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg		Purity:99.96%Clinical Data:LaunchedSize:10 mg, 50 mg	. 1975
Angiotensin II human acetate		Angiotensin II human TFA	
(Angiotensin II acetate; Ang II acetate; DRVYIHPF acetate)	Cat. No.: HY-13948A	(Angiotensin II TFA; Ang II TFA; DRVYIHPF TFA)	Cat. No.: HY-13948B
Angiotensin II human (Angiotensin II) acetate is a vasoconstrictor and a major bioactive peptide of the renin/angiotensin system.		Angiotensin II human (Angiotensin II) TFA is a vasoconstrictor and a major bioactive peptide of the renin/angiotensin system.	
Purity: 99.19% Clinical Data: Launched Size: 10 mM × 1 mL, 10 mg, 50 mg	A ou	Purity:99.49%Clinical Data:No Development ReportedSize:10 mg, 50 mg	r_{I}^{γ} on

Tel: 609-228-6898 Fax: 609-228-5909 Email: sales@MedChemExpress.com



Antineoplaston A10 Antitumor agent-36 Cat. No.: HY-128553 Cat. No.: HY-145288 Antineoplaston A10, a naturally occurring Antitumor agent-36 possesses potent substance in human body, is a Ras inhibitor anti-proliferative and anti-metastasis activities. potentially for the treatment of glioma, lymphoma, Antitumor agent-36 induces serious DNA damage and further leads to high expression of y-H2AX and astrocytoma and breast cancer. p53. H-N Purity: 98 58% >98% Purity: Clinical Data: Phase 2 Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg Size: 1 mg, 5 mg Antitumor agent-37 Antitumor agent-41 Cat. No.: HY-145289 Cat. No.: HY-144125 Antitumor agent-37 possesses potent Antitumor agent-41 (compound N-12), a potent anti-proliferative and anti-metastasis activities. antitumor agent, enhibits excellent antimigration Antitumor agent-37 induces serious DNA damage and and anti-invasion activity. Antitumor agent-41 further leads to high expression of y-H2AX and (compound N-12) induces tumor inhibition via tumor p53. necrosis and inflammatory response. H2N OF Purity: > 98% **Purity:** >98% Clinical Data: No Development Reported Clinical Data: No Development Reported Size: 1 mg, 5 mg Size: 1 mg, 5 mg Antitumor agent-42 Antitumor agent-43 Cat. No.: HY-144331 Cat. No.: HY-144340 Antitumor agent-42 (Compound 15h) is a Antitumor agent-43 (Compound 4B) is a potent N^{*}O bifunctional agent exhibiting both tubulin antitumor agent, with an IC_{50} of 0.5 μ M for (T-24 polymerized inhibition and NO-releasing cell). Antitumor agent-43 (Compound 4B) induces activities, resulting in potent anti-angiogenesis, cell cycle arrest at G2/M phase. colony formation inhibition, cell cycle arrest and apoptosis induction effects. Purity: > 98% Purity: >98% Clinical Data: No Development Reported Clinical Data: No Development Reported Size: 1 mg, 5 mg Size 1 mg, 5 mg Antitumor agent-44 Antitumor agent-45 Cat. No.: HY-144361 Cat. No.: HY-144394 Antitumor agent-44 (Compound 5n) disrupts the Antitumor agent-45 (Compound 21) could induce and mitochondrial homeostasis, induces cell cycle stimulate A549 cells apoptosis in G0/G1 and G2/M phase. Antitumor agent-45 (Compound 21) inhibits arrest and apoptosis in human adenocarcinoma cells. Antitumor agent-44 (Compound 5n) possesses c-Met expression to regulate the growth of tumor good anti-tumor activity in a lung-cancer-cell cells xenograft mice model. Purity: >98% **Purity:** >98% Clinical Data: No Development Reported Clinical Data: No Development Reported Size 1 mg, 5 mg Size 1 mg, 5 mg Antitumor agent-53 Antitumor agent-55 Cat. No.: HY-146743 Cat. No.: HY-146038 Antitumor agent-53 is a potent antitumor agent. Antitumor agent-55 (compound 5g) is a potent antitumor agent. Antitumor agent-55 effectively Antitumor agent-53 induces cell cycle arrest at the G2/M phase. Antitumor agent-53 inhibits the inhibits PC3, with an IC so of 0.91 µM. Antitumor agent-55 effectively inhibits the colony PI3K/AKT pathway to induce the apoptosis of HGC-27 cells. Antitumor agent-53 has the potential for formation, suppresses the cell migration in PC3. the research of gastrointestinal tumors. Purity: >98% >98% Purity: Clinical Data: No Development Reported Clinical Data: No Development Reported Size: 1 mg, 5 mg Size: 1 mg, 5 mg

AOH1160	Cat No : HY-120836	APG-1387	Cat No: HV-125593
AOH1160 is a potent, first-in-class, orally available small molecule proliferating cell nuclear antigen (PCNA) inhibitor, interferes with DNA replication, blocks homologous recombination-mediated DNA repair, causes cell-cycle arrest and induces apoptosis . Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		APG-1387, a bivalent SMAC mimetic and an IAP antagonist, blocks the activity of IAPs family proteins (XIAP, cIAP-1, cIAP-2, and ML-IAP). APG-1387 induces degradation of cIAP-1 and XIAP proteins, as well as caspase-3 activation and PARP cleavage, which leads to apoptosis. Purity: 99.46% Clinical Data: Phase 2 Size: 1 mg, 5 mg, 10 mg	
Aphidicolin	Cat. No.: HY-N6733	API-1	Cat. No.: HY-110077
Aphidicolin is an inhibitor of DNA polymerase α and δ , prevents mitotic cell division byinterfering with the activity of DNA polymerase.Aphidicolin is an antibiotic produced by the moldCephalosporium aphidicola.Purity: \geq 98.0%Clinical Data:No Development ReportedSize:1 mg	HOH HOH H	$\begin{array}{llllllllllllllllllllllllllllllllllll$	
Apitolisib	C + N - 10/ 122/C	Apocynin	
$\label{eq:GDC-0980; GNE 390; RG 7422)} \end{tabular} \begin{tabular}{lllllllllllllllllllllllllllllllllll$	Cat. No.: HY-13246	Apocynin is a selective NADPH-oxidase inhibitor with an IC ₅₀ of 10 μM. Purity: 99.95% Clinical Data: Phase 1 Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g	
Apogossypolone (ApoG2)	Cat. No.: HY-19551	Apoptolidin	Cat. No .: HY-126679
Apogossypolone (ApoG2) is an orally active Bcl-2 family proteins inhibitor with K ₁ values of 35, 25 and 660 nM for Bcl-2, Mcl-1 and Bcl-X ₁ , respectively. Apogossypolone shows antitumor activities, induces cell apoptosis and autophagy . Apogossypolone also has antifungal activity. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	Ha Ho Ho U O HO	Apoptolidin is a polyketide isolated from Nocardiopsis bacteria. Apoptolidin is a selective mitochondrial F_1F_0 ATPase inhibitor. Purity: $\geq 95.0\%$ Clinical Data: No Development Reported Size: 100 µg	
Apoptosis inducer 4	Cat. No. : HY-146092	Apoptozole (Apoptosis Activator VII)	Cat. No.: HY-15098
Apoptosis inducer 4 (Compound 12b) is an apoptosis inducer with anticancer activities.	mongh	Apoptozole (Apoptosis Activator VII) is an inhibitor of the ATPase domain of Hsc70 and Hsp70, with K_ds of 0.21 and 0.14 μ M, respectively, and can induce apoptosis.	FL LF P LF NN NC NHG
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:99.81%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg	

Apremilast		Apremilast-d5	
(CC-10004)	Cat. No.: HY-12085	(CC-10004-d5)	Cat. No.: HY-12085S
Apremilast (CC-10004) is an orally available inhibitor of type-4 cyclic nucleotide phosphodiesterase (PDE-4) with an IC ₅₀ of 74 nM. Apremilast inhibits TNF-α release by lipopolysaccharide (LPS) with an IC ₅₀ of 104 nM. Purity: 99.87% Clinical Data: Launched Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	$ \begin{array}{c} $	Apremilast D5 (CC-10004 D5) is a deuterium labeledApremilast. Apremilast is an orally availableinhibitor of type-4 cyclic nucleotidephosphodiesterase (PDE-4) with an IC ₅₀ of 74 nM.Apremilast inhibits TNF- α release bylipopolysaccharide (LPS) with an IC ₅₀ of 104 nM.Purity:>98%Clinical Data:No Development ReportedSize:1 mg	
APTO-253		AQX-435	
(LOR-253; LT-253)	Cat. No.: HY-16291		Cat. No.: HY-136268
APTO-253 (LOR-253) is a small molecule that inhibits c-Myc expression, stabilizes G-quadruplex DNA, and induces cell cycle arrest and apoptosis in acute myeloid leukemia cells.		AQX-435 is a potent SHIP1 phosphatase activator. AQX-435 reduces PI3K activation downstream of the B-cell receptor (BCR) and induces apoptosis of malignant B cells, and reduces lymphoma growth.	HO U
Clinical Data: Phase 1 Size: 10 mM x 1 mL 5 mg 10 mg 50 mg 100 mg		Clinical Data: No Development Reported	
512e. 10 million × 1 mill, 5 million, 10 million, 100 million		Size. 1 mg, 5 mg	
ar Turmoropo		ar Turmarana d2	
((+)-ar-Turmerone)	Cat No HV-N6703	((+)-ar-Turmerone-d3)	Cat No : HV-N6703S
ar-Turmerone ((+)-ar-Turmerone) is a major bioactive compound of the herb Curcuma longa with anti-tumorigenesis and anti-inflammatory activities. ar-Turmerone activates apoptotic protein in human lymphoma U937 cells. Purity: >98% Clinical Data: No Development Reported Size: 5 mg, 10 mg		ar-Turmerone-d3 ((+)-ar-Turmerone-d3) is the deuterium labeled ar-Turmerone. ar-Turmerone ((+)-ar-Turmerone) is a major bioactive compound of the herb Curcuma longa with anti-tumorigenesis and anti-inflammatory activities. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	
Arctigenin		ARD-61	
((-)-Arctigenin)	Cat. No.: HY-N0035		Cat. No.: HY-139659
Arctigenin ((-)-Arctigenin), a biologically active lignan, can be used as an antitumor agent. Arctigenin exhibits potent antioxidant, anti-inflammatory and antiviral (influenza A virus) activities.	о Ср. А. С. С.	ARD-61 is a highly potent, effective and specific PROTAC androgen receptor (AR) degrader. ARD-61 potently and effectively induces AR and progesterone receptors (PR) degradation in AR+ cancer cell lines.	- Altrono ostar
Purity: 99.69%		Purity: >98%	
Clinical Data: Phase 2		Clinical Data: No Development Reported	
SIZE: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg		Size: 1 mg, 5 mg	
Ardisiacrispin B		Aristolactam I (Aristololactam: Aristolactam)	Cat No. HV-NI2013
Ardisiacrispin B displays cytotoxic effects in	Cat. No.: 111-110150	Aristololactam I (AL-I), is the main metabolite of	_/~Q
multi-factorial drug resistant cancer cells via ferroptotic and apoptotic cell death.	An in the	aristolochic acid I (AA-I), participates in the processes that lead to renal damage.	
			NH NH
Purity: >98%		Purity: 99.69%	_0
Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg		Clinical Data: No Development Reported Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg	
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ASK1-IN-2	Cat. No.: HY-131969	ASK1-IN-3	Cat. No.: HY-146729
ASK1-IN-2 is a potent and orally active inhibitor of apoptosis signal-regulating kinase 1 (ASK1) , with an IC_{so} of 32.8 nM. ASK1-IN-2 can be used for the research of ulcerative colitis.		ASK1-IN-3 is a potent and selective ASK1 kinase inhibitor with IC_{s0} of 33.8 nM, as well as inhibits several cell cycle regulating kinases. ASK1-IN-3 has strong HepG2 cancer cells apoptosis induction and potent cell cycle arrest activities.	
Purity:98.49%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	00 mg	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	Ю
ASP3026	Cat No: HV-13326	Asperosaponin VI	Cat No. HV-N0265
ASP3026 is a potent, selective and orally active inhibitor of anaplastic lymphoma kinase (ALK) . ASP3026 induces apoptosis of tumor cells. ASP3026 can be used for the research of non-small cell lung cancer (NSCLC).		Asperosaponin VI, A saponin component from Dipsacus asper wall, induces osteoblast differentiation through BMP2/p38 and ERK1/2 pathway.	
Purity: 99.90% Clinical Data: Phase 1 Size: 10 mM × 1 mL, 50 mg, 100 mg		Purity:98.73%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 20 mg	Он он
		477510	
Astragalin (Astragaline; 3-Glucosylkaempferol; Kaempfe 3-B-D-glucopyranoside)	cat. No.: HY-N0015	AT 7519 (AT7519M)	Cat. No.: HY-50940
Astragalin (kaempferol-3-O-glucoside) is a flavonoid with anti-inflammatory activity and newly found in persimmon leaves and green tea seeds.	HO-CHOHOHOHOHOHOHOHOHOHOHOHOHOHOHOHOHOHO	AT7519 (AT7519M) as a potent inhibitor of CDKs , with IC ₅₀ s of 210, 47, 100, 13, 170, and <10 nM for CDK1, CDK2, CDK4 to CDK6, and CDK9, respectively.	
Purity:99.85%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg	тон	Purity: 99.76% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	н
AT7519 Hydrochloride	Cat. No. : HY-50943	AT9283	Cat. No.: HY-50514
AT7519 Hydrochloride is a potent inhibitor of CDKs, with IC_{so} s of 210, 47, 100, 13, 170, and <10 nM for CDK1, CDK2, CDK4 to CDK6, and CDK9, respectively.		AT9283 is a multi-targeted kinase inhibitor with potent activity against Aurora A/B, JAK2/3, Abl (T315I) and Flt3 (IC_{so} s ranging from 1 to 30 nM). AT9283 inhibits growth and survival of multiple solid tumors in vitro and in vivo.	
Purity: 99.29% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	H-CI	Purity: 99.70% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 50 mg, 100) mg
Atezolizumah			
(MPDL3280A)	Cat. No.: HY-P9904		Cat. No.: HY-15003
Atezolizumab (MPDL3280A) is a selective humanized monoclonal IgG1 antibody against programmed death ligand 1 (PD-L1), used for cancer research.	Atezolizumab	ATH686 is a potent, selective and ATP-competitive FLT3 inhibitor. ATH686 target mutant FLT3 protein kinase activity and inhibit the proliferation of cells harboring FLT3 mutants via induction of apoptosis and cell cycle inhibition. ATH686 has antileukemic effects.	A A A A A A A A A A A A A A A A A A A
Purity: 98.98% Clinical Data: Launched Size: 1 mg, 5 mg, 25 mg, 50 mg		Purity:99.58%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg	



47.628		47960	
R2 020	Cat. No.: HY-11004	ALSOO	Cat. No.: HY-10411
AZ 628 is a pan-Raf kinase inhibitor with IC _{so} s of 105, 34 and 29 nM for B-Raf, B-RafV600E , and c-Raf-1 , respectively.	in the second	AZ960 is a potent and specific inhibitor of the JAK2 kinase with a K_i of 0.45 nM.	
Purity:99.86%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg		Purity:97.15%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 50 mg, 100) mg
AZA1		Azadirachtin	
(Rac1/Cdc42-IN-1)	Cat. No.: HY-136383		Cat. No.: HY-126741
AZA1 is a potent dual inhibitor of Rac1 and Cdc42 . AZA1 induces prostate cancer cells apoptosis and inhibits prostate cancer cells proliferation, migration and invasion.		Azadirachtin, one of the most promising botanical insecticides, is widely used for pest control. Azadirachtin induces apoptosis in insect cell lines, including Sf9, SL-1 and BTI-Tn-5B1-4.	Hand to the second seco
Purity: 98.65% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity:98.05%Clinical Data:No Development ReportedSize:1 mg, 5 mg	0
Azathioprine		AZD-3463	
(BW 57-322)	Cat. No.: HY-B0256	(ALK/IGF1R inhibitor)	Cat. No.: HY-15609
Azathioprine(Azasan, Imuran; BW 57-322) is a drug that suppresses the immune system and is used in organ transplantation and autoimmune disease. Target: Azathioprine is an immunosuppressive antimetabolite pro-drug. Purity: 99.98%		AZD-3463 (ALK/IGF1R inhibitor) is an orally active ALK/IGF1R inhibitor, with a K _i of 0.75 nM for ALK. AZD3463 induces apoptosis and autophagy in neuroblastoma cells. Purity: 99.96%	
Clinical Data: Launched Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g		Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
AZD-8055	Cat. No.: HY-10422	AZD0156	Cat. No.: HY-100016
AZD-8055 is a potent, selective, and orally bioavailable ATP-competitive mTOR kinase inhibitor with an IC_{50} of 0.8 nM. AZD-8055 inhibits both mTORC1 and mTORC2.		AZD0156 is a potent, selective and orally active ATM inhibitor with an IC_{so} of 0.58 nM. AZD0156 inhibits the ATM-mediated signaling, prevents DNA damage checkpoint activation, disrupts DNA damage repair, and induces tumor cell apoptosis .	installer.
Purity: 99.60% Clinical Data: Phase 1 Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg		Purity: 99.82% Clinical Data: Phase 1 Size: 5 mg, 10 mg, 50 mg, 100 mg	
A7D0424		A7D1208	
	Cat. No.: HY-112314		Cat. No.: HY-15604
AZD0424 is an orally active, and dual selective Src/Abl kinase inhibitor with potential antineoplastic activity. AZD0424 induces apoptosis and cell cycle arrest in lymphoma cells.		AZD1208 is an orally bioavailable, highly selective PIM kinases inhibitor.	O S NH2
Purity:>98%Clinical Data:Phase 1Size:1 mg, 5 mg		Purity: 99.90% Clinical Data: Phase 1 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 2	200 mg

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AZD1208 hydrochloride		AZD5582	
	Cat. No.: HY-15604A		Cat. No.: HY-12600
AZD1208 hydrochloride is an orally bioavailable, highly selective PIM kinases inhibitor.		AZD5582 is an antagonist of the inhibitor of apoptosis proteins (IAPS) , which binds to the BIR3 domains cIAP1, cIAP2, and XIAP with IC ₅₀ s of 15, 21, and 15 nM, respectively. AZD5582 induces apoptosis .	*****
Purity: >98% Clinical Data: Phase 1 Size: 1 mg, 5 mg	H-Ci	Purity: 98.11% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg	
AZD5582 dihydrochloride	Cat. No.: HY-110346	Azoramide	Cat. No.: HY-18705
AZD5582 dihydrochloride is an antagonist of the inhibitor of apoptosis proteins (IAPs) , which binds to the BIR3 domains cIAP1, cIAP2, and XIAP with IC_{so} of 15, 21, and 15 nM, respectively. AZD5582 induces apoptosis .	ulity	Azoramide is a small-molecule modulator of the unfolded protein response with antidiabetic activity. in vitro: Azoramide is a dual-function endoplasmic reticulum (ER) modulator.	~~ ^{\$} N~ ^{\$} ~O~a
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity: 98.63% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	L00 mg
Azoxystrobin	Cat. No. : HY-B0849	Azoxystrobin-d3	Cat. No.: HY-B0849S1
Azoxystrobin is a broad-spectrum β-methoxyacrylate fungicide. Azoxystrobin inhibits mitochondrial respiration by binding to the Qo site of the cytochrome bc1 complex and inhibiting electron transfer. Purity: 99.06%		Azoxystrobin-d3 is deuterium labeled Azoxystrobin. Azoxystrobin is a broad-spectrum β-methoxyacrylate fungicide. Azoxystrobin inhibits mitochondrial respiration by binding to the Qo site of the cytochrome bc1 complex and inhibiting electron transfer. Purity: >98% Clicical Data Na Davalanzara Davata	
Size: 10 mM × 1 mL, 50 mg		Size: 1 mg, 5 mg	
Azoxystrobin-d4	Cat. No.: HY-B0849S	AZT triphosphate (3'-Azido-3'-deoxythymidine-5'-triphosphate)	Cat. No. : HY-116364
Azoxystrobin-d4 is deuterium labeled Azoxystrobin. Azoxystrobin is a broad-spectrum β-methoxyacrylate fungicide. Azoxystrobin inhibits mitochondrial respiration by binding to the Qo site of the cytochrome bc1 complex and inhibiting electron transfer.		AZT triphosphate (3'-Azido-3'-deoxythymidine-5'-triphosphate) is a active triphosphate metabolite of Zidovudine (AZT). AZT triphosphate exhibits antiretroviral activity and inhibits replication of HIV .	John Charles
Purity:>98%Clinical Data:Size:1 mg, 5 mg		Purity: >98% Clinical Data: No Development Reported Size: 1 mg	
AZT triphosphate TEA (3'-Azido-3'-deoxythymidine-5'-triphosphate TEA)	Cat. No. : HY-116364A	B-355252	Cat. No.: HY-120553
AZT triphosphate TFA (3'-Azido-3'-deoxythymidine-5'-triphosphate TFA) is a active triphosphate metabolite of Zidovudine (AZT). AZT triphosphate TFA exhibits antiretroviral activity and inhibits replication of HIV .		B355252, a phenoxy thiophene sulfonamide small molecule, is a potent NGF receptor agonist. B355252 potentiates NGF-induced neurite outgrowth.	
Purity: >98% Clinical Data: No Development Reported Size: 1 mg		Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	00



Batabulin		Batabulin sodium	
(T138067)	Cat. No : HY-13563	(T138067 sodium)	Cat. No : HY-135634
Batabulin (T138067) is an antitumor agent, which binds covalently and selectively to a subset of the β-tubulin isotypes, thereby disrupting microtubule polymerization. Batabulin affects cell morphology and leads to cell-cycle arrest ultimately induces apoptotic cell death. Purity: 99.91% Clinical Data: Phase 3 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg		Batabulin sodium (T138067 sodium) is an antitumor agent, which binds covalently and selectively to a subset of the β-tubulin isotypes, thereby disrupting microtubule polymerization. Batabulin sodium affects cell morphology and leads to cell-cycle arrest ultimately induces apoptotic cell death. Purity: 99.79% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg	
Bax activator-1	Cat. No.: HY-122760	Bax BH3 peptide (55-74), wild type	Cat. No.: HY-P2466
Bax activator-1 (compound 106) is a Bax activator that induces Bax-dependent tumor cell apoptosis.	Dun Junger	Bax BH3 peptide (55-74), wild type is a 20-amino acid Bax BH3 peptide (Bax 1) capable of inducing apoptosis in a variety of cell line models.	STKKLSECLKRIGDELDSNM
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
BAY 11-7082		BAY 11-7085	
(BAY 11-7821)	Cat. No.: HY-13453	(BAY 11-7083)	Cat. No.: HY-10257
BAY 11-7082 is an IxB α phosphorylation and NF- κ B inhibitor. BAY 11-7082 selectively and irreversibly inhibits the TNF- α -induced phosphorylation of I κ B- α , and decreases NF- κ B and expression of adhesion molecules.	C o o o o o o o o o o o o o o o o o o o	BAY 11-7085 (BAY 11-7083) is an inhibitor of NF-κB activation and phosphorylation of IκBα; it stabilizes IκBα with an IC _{so} of 10 μM.	× o o
Purity:99.98%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity: 99.99% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	.00 mg
BAY 61-3606		BAY 61-3606 dihydrochloride	
	Cat. No.: HY-76474		Cat. No.: HY-14985
BAY 61-3606 is an orally available, ATP-competitive, reversible and highly selective Syk inhibitor with a K_i of 7.5 nM and an IC ₅₀ of 10 nM. BAY 61-3606 reduces ERK1/2 and Akt phosphorylation in neuroblastoma cell. Purity: 98.21% Clinical Data: No Development Reported Size: 1 mg, 5 mg		$\begin{array}{llllllllllllllllllllllllllllllllllll$	
RAV1082/439		RC-1258	
5.11002435	Cat. No.: HY-100886	50 1250	Cat. No.: HY-129087
BAY1082439 is an orally bioavailable, selective PI3K $\alpha/\beta/\delta$ inhibitor. BAY1082439 also inhibits mutated forms of PIK3CA. BAY1082439 is highly effective in inhibiting Pten-null prostate cancer growth.	PH O O N N N N N N N N N N N N N N	BC-1258, an F-box/LRR-repeat protein 2 (FBXL2) activator, can stabilize and upregulate FBXL2 levels. BC-1258 induces apoptosis of tumorigenic cells, and profoundly inhibits tumor formation in mice.	agentas
Purity:99.46%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	<u>o</u>	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	

Bcl-2-IN-8	Cat. No.: HY-144819	Bendamustine (SDX-105 free base)	Cat. No. : HY-13567
Bcl-2-IN-8 is a potent anticancer agent.Bcl-2-IN-8 shows anti-proliferative activityagainst both drug-sensitive and drug-resistantcancer cells. Bcl-2-IN-8 induce apoptosis and cellcycle arrest at G1 phase. Bcl-2-IN-8 inhibits cellmigration in a dose-dependent manner.Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	Jon Martin	Bendamustine (SDX-105 free base), a purine analogue, is a DNA cross-linking agent. Bendamustine activates DNA-damage stress response and apoptosis. Bendamustine has potent alkylating, anticancer and antimetabolite properties. Purity: ≥98.0% Clinical Data: Launched Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg	CI~~N~CI
Bendamustine hydrochloride (SDX-105)	Cat. No.: HY-B0077	Bendamustine-d4 hydrochloride	Cat. No. : HY-B0077S
Bendamustine hydrochloride (SDX-105), a purine analogue, is a DNA cross-linking agent. Bendamustine hydrochloride activats DNA-damage stress response and apoptosis. Bendamustine hydrochloride has potent alkylating, anticancer and antimetabolite properties. Purity: 98.94% Clinical Data: Launched Size: 10 mM × 1 mL, 25 mg, 100 mg, 200 mg, 500	асти стран на mg	Bendamustine-d4 hydrochloride is the deuterium labeled Bendamustine hydrochloride. Bendamustine hydrochloride (SDX-105), a purine analogue, is a DNA cross-linking agent. Bendamustine hydrochloride activats DNA-damage stress response and apoptosis. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	a ~ N ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~
		D. THE	
Bendamustine-d8 hydrochloride		Benidipine	
(SDX-105-d8)	Cat. No.: HY-B0077S1	(KW-3049 free base)	Cat. No.: HY-B1448A
Bendamustine-d8 (hydrochloride) is deuterium labeled Bendamustine (hydrochloride). Bendamustine hydrochloride (SDX-105), a purine analogue, is a DNA cross-linking agent. Bendamustine hydrochloride activats DNA-damage stress response and apoptosis. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Benidipine is a potent and orally active calcium channel antagonist. Benidipine shows anti-apoptosis effects in ischaemic/reperfused myocardial cells. Benidipine increases the activity of endothelial cell-type nitric oxide synthase and improves coronary circulation in hypertensive rats. Purity: >98% Clinical Data: Launched Size: 1 mg, 5 mg	of the Carl
Benzbromarone	Cat No: HY-B1135	Benzbromarone-d5	Cat No: HY-B1135S
Benzbromarone is a highly effective and well tolerated non-competitive inhibitor of xanthine oxidase, used as an uricosuric agent, used in the treatment of gout.	C+C+C+Br O+C+C+OH	Benzbromarone-d5 is deuterium labeled Benzbromarone.	
Purity:99.80%Clinical Data:LaunchedSize:10 mM × 1 mL, 100 mg	ы	Purity:>98%Clinical Data:Size:1 mg, 5 mg	Br
Benzoylpaeoniflorin	Cat. No.: HY-N0852	Benzyl isothiocyanate	Cat. No.: HY-77813
Benzoylpaeoniflorin, a natural product from Chinese paeony root, has the potential for coronary heart disease by decreasing apoptosis.	HO OF OF	Benzyl isothiocyanate is a member of natural isothiocyanates with antimicrobial activity. Benzyl isothiocyanate potent inhibits cell mobility, migration and invasion nature and matrix metalloproteinase-2 (MMP-2) activity of murine melanoma cells.	s ^{zczN}
Purity: ≥99.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	Û.,	Purity: ≥98.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 100 mg	

Benzyl isothiocyanate-d7	Berbamine dihydrochloride
Benzyl isothiocyanate-d7 is the deuterium labeled Benzyl isothiocyanate. Benzyl isothiocyanate is a member of natural isothiocyanates with antimicrobial activity.	Cat. NO: HY-NO/14A Berbamine dihydrochloride is an inhibitor of NF-κB activity with remarkable anti-myeloma efficacy.
Purity: >98% Clinical Data: No Development Reported Size: 5 mg, 50 mg	Purity: 96.62% Clinical Data: Launched Size: 10 mM × 1 mL, 200 mg, 500 mg
Bergenin (Cuscutin) Cat. No.: HY-N0017	beta-Mangostin (β-Mangostin) Cat. No.: HY-N0941
Bergenin is a cytoprotective and antioxidative polyphenol found in many medicinal plants. Bergenin has a wide spectrum activities such as hepatoprotective, antiinflammatory, immunomodulatory, antitumor, antiviral, and antifungal properties. Purity: 99.63% Clinical Data: Launched Size: 10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 50 mg	beta-Mangostin (β -Mangostin) is a xanthone compound present in Cratoxylum arborescens, with antibacterial and antimalarial activities. beta-Mangostin exhibits antimycobacterial activity against Mycobacterium tuberculosis with an MIC of 6.25 µg/mL. Purity: 99.74% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg
Beta-Sitosterol (purity>75%) (β-Sitosterol (purity>75%); 22,23-Dihydrostigmasterol (purity>75%)) Cat. No.: HY-N0171B	Beta-Sitosterol (purity>80%)(β-Sitosterol (purity>80%);22,23-Dihydrostigmasterol (purity>80%))Cat. No.: HY-N0171
Beta-Sitosterol (purity>75%) includes 75% β -sitosterol and 10% campesterol. Beta-Sitosterol is a plant sterol.	Beta-Sitosterol (purity>80%) includes β -sitosterol (>80%), stigmasterol, campesterol and brassicasterol mainly. Beta-Sitosterol is a plant sterol.
Purity: ≥95.0% (purity>75%) Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg	Purity: ≥80.0% (purity>80%) Clinical Data: No Development Reported Size: 100 mg, 1 g, 5 g
Beta-Sitosterol (purity>98%) (β-Sitosterol (purity>98%); 22,23-Dihydrostigmasterol (purity>98%)) Cat. No.: HY-N0171A	Beta-Zearalanol Cat. No.: HY-N6740
Beta-Sitosterol (purity>98%) is a plant sterol. Beta-Sitosterol (purity>98%) interfere with multiple cell signaling pathways, including cell cycle, apoptosis, proliferation, survival, invasion, angiogenesis, metastasis and inflammation.	Beta-Zearalenol is an mycotoxin produced by Fusarium spp, which causes apoptosis and oxidative stress in mammalian reproductive cells. Beta-Zearalenol is the derivative of zearalenone (ZEA) which can conjugate with glucuronic acid.
Purity: ≥98.0% (purity=98%) Clinical Data: No Development Reported Size: 10 mg, 25 mg, 50 mg, 100 mg	Purity: >98% Clinical Data: No Development Reported Size: 5 mg, 10 mg
Betamethasone Cat. No.: HY-13570	Betamethasone hydrochloride Cat. No.: HY-13570A
Betamethasone is a synthetic glucocorticoid with anti-inflammatory and immunosuppressive activities. Betamethasone accelerates fetal lung maturation and induces gene expression and apoptosis.	Betamethasone hydrochloride is a synthetic glucocorticoid with anti-inflammatory and immunosuppressive activities. Betamethasone hydrochloride accelerates fetal lung maturation and induces gene expression and apoptosis. How the hydrochloride accelerates fetal lung maturation and induces gene expression and apoptosis.
Purity: 99.97% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 500 mg	Purity: >98% HCI Clinical Data: Launched Size: 1 mg, 5 mg

Betamethasone-d5	Cat. No.: HY-13570S	Betamethasone-d5-1	Cat. No.: HY-13570S1
Betamethasone-d5 is the deuterium labeledBetamethasone. Betamethasone is a syntheticglucocorticoid with anti-inflammatory andimmunosuppressive activities. Betamethasoneaccelerates fetal lung maturation and induces geneexpression and apoptosis.Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Betamethasone-d5-1 is deuterium labeledBetamethasone. Betamethasone is a syntheticglucocorticoid with anti-inflammatory andimmunosuppressive activities. Betamethasoneaccelerates fetal lung maturation and induces geneexpression and apoptosis.Purity:>98%Clinical Data:Size:1 mg, 5 mg	
BETd-260 (ZBC 260)	Cat. No.: HY-101519	Betulin (Trochol)	Cat. No.: HY-N0083
BETd-260 (ZBC 260) is a PROTAC connected by ligands for Cereblon and BET , with as low as 30 pM against BRD4 protein in R54;11 leukemia cell line. BETd-260 potently suppresses cell viability and robustly induces apoptosis in hepatocellular carcinoma (HCC) cells.	2007	Betulin (Trochol), is a sterol regulatory element-binding protein (SREBP) inhibitor with an IC ₅₀ of 14.5 μM in K562 cell line.	
Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg, 50 mg		Clinical Data: No Development Reported Size: 50 mg, 100 mg, 200 mg	
Betulinic acid		BG45	
Betulinic acid; Betulic acid) Betulinic acid is a natural pentacyclic triterpenoid, acts as a eukaryotic topoisomerase I inhibitor, with an IC _{so} of 5 μM, and possesses anti-HIV, anti-malarial, anti-inflammatory and anti-tumor properties.		BG45 is an HDAC class I inhibitor with selectivity for HDAC3 (IC50 = 289 nM). It inhibits HDAC1, HDAC2, and HDAC6 with greatly reduced potency (IC50s = 2, 2.2, and >20 μ M, respectively).	HY-18/12
Purity: ≥98.0% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 100 mg, 200 mg, 500 mg		Purity: 99.95% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
BGT226 maleate (NVP-BGT226 maleate)	Cat. No.: HY-13334	BH3 hydrochloride	Cat. No.: HY-P2343
BGT226 (NVP-BGT226 maleate) is a PI3K (with IC _{so} s of 4 nM, 63 nM and 38 nM for PI3K α , PI3K β and PI3K γ) / mTOR dual inhibitor which displays potent growth-inhibitory activity against human head and neck cancer cells.	N C C C C C C C C C C C C C C C C C C C	BH3 hydrochloride, a BBB penetrated peptide, provoke apoptosis either by direct activation of pro-apoptotic Bax/Bak or by neutralizing anti-apoptotic Bcl-2 proteins (Bcl-2, Bcl-XL, Bcl-w, Mcl-1 and A-1) via their BH3 domian.	WIAGEURRIGDEFNAYYARR (HCI sat)
Purity: 99.73% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	
BI 2536	Cat. No.: HY-50698	BI-6C9	Cat. No.: HY-103661
BI 2536 is a dual PLK1 and BRD4 inhibitor with IC_{50} s of 0.83 and 25 nM, respectively. BI-2536 suppresses IFNB (encoding IFN-β) gene transcription.		BI-6C9 is a highly specific BH3 interacting domain (Bid) inhibitor, which prevents mitochondrial outer membrane potential (MOMP) and mitochondrial fission, and protects the cells from mitochondrial apoptosis inducing factor (AIF) release and caspase-independent cell death in neurons.	~0, ⁴ ~, ¹ 0,0,"
Purity: 99.95% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 5 mg, 25 mg, 50 mg, 100 mg	Ų _N .	Purity:98.24%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg	

BI-847325	Cat. No.: HY-18955	Biatractylolide ((+)-Biatractylolide)	Cat. No.: HY-N10131
BI-847325 is an ATP competitive dual inhibitor of MEK and aurora kinases (AK) with IC_{50} values of 4 and 15 nM for human MEK2 and AK-C, respectively.	and an	Biatractylolide is a compound isolated from the ethyl acetate extract of Atractylodes macrocephala. Biatractylolide has antitumor and antioxidant activities.	
Purity: 98.66% Clinical Data: Phase 1 Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	مر 0 mg	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	Ŷ₩ Ħ
BIBR 1532	Cat. No. : HY-17353	Bigelovin	Cat. No. : HY-116506
BIBR 1532 is a potent, selective and non-competitive telomerase inhibitor with IC_{50} of 100 nM in a cell-free assay.	COLO HOLO	Bigelovin, a sesquiterpene lactone isolated from Inula helianthus-aquatica, is a selective retinoid X receptor α agonist. Bigelovin suppresses tumor growth through inducing apoptosis and autophagy via the inhibition of mTOR pathway regulated by ROS generation.	H H H H H H H H H H H H H H H H H H H
Purity:99.94%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg, 50 mg, 100 mg		Purity:99.81%Clinical Data:No Development ReportedSize:5 mg, 10 mg	0
Bilobalide		BIO-acetoxime	
Bilobalide, a sesquiterpene trilactone constituent of Ginkgo biloba, inhibits the NMDA-induced efflux of choline with an IC_{so} value of 2.3 μ M. Bilobalide prevents apoptosis through activation of the PI3K/Akt pathway in SH-SY5Y cells. Exerts protective and trophic effects on neurons.		BIO-acetoxime (BIA) is a potent and selective GSK-3 inhibitor, with $IC_{so}s$ of both 10 nM for GSK-3 α/β . BIO-acetoxime has anticonvulsant and anti-infection activity.	
Purity: ≥98.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg	н	Purity: ≥98.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	00 mg
Diovrnifi		PiD inducer V	
(DR5 Activator)	Cat. No.: HY-18377		Cat. No.: HY-110188
Bioymifi (DR5 Activator), a potent TRAIL receptor DR5 activator, binds to the extracellular domain (ECD) of DR5 with a K_d of 1.2 μ M. Bioymifi can act as a single agent to induce DR5 clustering and aggregation, leading to apoptosis.	R-O-J-S-O-J-O-J-HI	BiP inducer X, a selective inducer of immunoglobulin heavy chain binding protein (BiP)/GRP78, is an effective ER (endoplasmic reticulum) stress inhibitor.	
Purity: ≥98.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	0 mg	Purity:99.88%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
Birinapant (TL32711)	Cat. No. : HY-16591	Bisdemethoxycurcumin (Curcumin III; Didemethoxycurcumin)	Cat. No. : HY-N0007
Birinapant (TL32711), a bivalent Smac mimetic, is a potent antagonist for XIAP and cIAP1 with K _d s of 45 nM and less than 1 nM, respectively.	and the second s	Bisdemethoxycurcumin(Curcumin III; Didemethoxycurcumin) is a natural derivative of curcumin with anti-inflammatory and anti-cancer activities.	ностение
Purity: 99.70% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity: ≥98.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg	

Bisdemethoxycurcumin-d8 (Curcumin III-d8; Didemethoxycurcumin-d8)	Cat. No.: HY-N0007S	Bisindolylmaleimide VIII acetate (Ro 31-7549 acetate; Bis VIII acetate)	Cat. No. : HY-129624A
Bisdemethoxycurcumin-d8 (Curcumin III-d8) is the deuterium labeled Bisdemethoxycurcumin. Bisdemethoxycurcumin(Curcumin III; Didemethoxycurcumin) is a natural derivative of curcumin with anti-inflammatory and anti-cancer activities.		BisindolyImaleimide VIII acetate (Ro 31-7549 acetate) is a potent and selective protein kinase C (PKC) inhibitor with an IC_{s0} of 158 nM for rat brain PKC.	CT N C NH
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:99.70%Clinical Data:No Development ReportedSize:5 mg	
BJE6-106 (B106)	Cat. No. : HY-117800	Bleomycin A5 hydrochloride (Pingyangmycin hydrochloride)	Cat. No.: HY-125918
BJE6-106 (B106) is a potent, selective 3^{rd} generation PKC δ inhibitor with an IC ₅₀ of 0.05 μ M and targets selectivity over classical PKC isozyme PKC α (IC ₅₀ =50 μ M). BJE6-106 (B106) induces caspase-dependent apoptosis . BJE6-106 (B106) possesses tumor-specific effect.		Bleomycin A5 (Pingyangmycin) hydrochloride is an anti-neoplastic glycoprotein antibiotic . Bleomycin A5 suppresses Drp1-mediated mitochondrial fission and induces apoptosis in human nasal polyp-derived fibroblasts.	Sill for
Purity: 98.17% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	0-0	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
BLM-IN-1	Cat. No.: HY-111756	BML-210	Cat. No.: HY-19350
BLM-IN-1 (compound 29) is an effective Bloom syndrome protein (BLM) inhibitor, with a strong BLM binding K_0 of 1.81 μ M and an IC ₅₀ of 0.95 μ M for BLM. Induces DNA damage response, as well as apoptosis and proliferation arrest in cancer cells.	mitta	BML-210 is a novel HDAC inhibitor, and its mechanism of action has not been characterized.	
Purity:99.08%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg		Purity: 96.38% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
BML-277		BMS-1	
(Chk2 Inhibitor II)	Cat. No.: HY-13946	(PD-1/PD-L1 inhibitor 1)	Cat. No.: HY-19991
BML-277 is a selective checkpoint kinase 2 (Chk2) inhibitor with an $\rm IC_{50}$ of 15 nM.	*** ¹ 077-0-0 ⁰	BMS-1 is an inhibitor of the PD-1/PD-L1 protein/protein interaction (IC ₅₀ between 6 and 100 nM).	ملەرىپ
Purity:98.49%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg, 50 mg		Purity:99.56%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	00 mg
BMS-202	Cat. No.: HY-19745	BMS-536924	Cat. No.: HY-10262
BMS-202 is a potent and nonpeptidic PD-1/PD-L1 complex inhibitor with an IC_{s0} of 18 nM and a $K_{\rm p}$ of 8 μ M. BMS-202 binds to PD-L1 and blocks human PD-1/PD-L1 interaction. BMS-202 has antitumor activity.	1 in the	BMS-536924 is an orally active, competitive and selective insulin-like growth factor receptor (IGF-1R) kinase and insulin receptor (IR) inhibitor with IC ₅₀ s of 100 nM and 73 nM, respectively. BMS-536924 has anti-cancer activity.	
Purity: 99.39% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	00 mg	Purity: 99.74% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	CI
BMS-833923 (XL-139)	Cat. No.: HY-13809	BO-264	Cat. No.: HY-135960
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BMS-833923 (XL-139) is an orally bioavailable small-molecule inhibitor of Smoothened with potential antineoplastic activity; inhibits BODIPY cyclopamine binding to SMO in a dose-dependent manner with an IC50 of 21 nM.	atta to t	BO-264 is a highly potent and orally active transforming acidic coiled-coil 3 (TACC3) inhibitor with an IC_{so} of 188 nM and a K_d of 1.5 nM. BO-264 specifically blocks the function of FGFR3-TACC3 fusion protein.	CN IN STORY
Purity: 98.21% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg		Purity:99.63%Clinical Data:No Development ReportedSize:10 mg, 50 mg, 100 mg, 250 mg	
BOC-D-FMK	Cat. No.: HY-13229	Bornyl acetate	Cat. No.: HY-N0756
Boc-D-FMK is a cell-permeable, irreversible and broad spectrum caspase inhibitor. Boc-D-FMK inhibits apoptosis stimulated by TNF- α with an IC _{so} of 39 μ M.	Xoly Co	Bornyl acetate is a potent odorant, exhibiting one of the highest flavor dilution factor (FD factor). Bornyl acetate possesses anti-cancer activity.	
Purity: ≥95.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 50 mg		Purity:≥97.0%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 100 mg, 500 mg	Relative Stereouternistry
Dowelidin		Berterenik	
(Treponemycin)	Cat. No.: HY-N6742	(PS-341; LDP-341; NSC 681239)	Cat. No.: HY-10227
Borrelidin (Treponemycin) is a bacterial and eukaryal threonyl-tRNA synthetase inhibitor which is a nitrile-containing macrolide antibiotic isolated from Streptomyces rochei. Borrelidin is an inhibitor of Cdc28/Cln2 of the budding yeast, with an IC ₅₀ of 24 μ M. Purity: \geq 98.0% Clinical Data No Daveloament Bacanted		Bortezomib (PS-341) is a reversible and selective proteasome inhibitor, and potently inhibits 20S proteasome ($K_{=}$ 0.6 nM) by targeting a threonine residue. Bortezomib disrupts the cell cycle, induces apoptosis, and inhibits NF- κ B.	
Size: 500 µg, 1 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 5	200 mg
Bortezomib-d8 (PS-341-d8; LDP-341-d8; NSC 681239-d8)	Cat. No. : HY-102275	bpV(phen)	Cat. No. : HY-136065
Bortezomib-d8 (PS-341-d8) is the deuterium labeled Bortezomib. Bortezomib (PS-341) is a reversible and selective proteasome inhibitor, and potently inhibits 20S proteasome (K _i =0.6 nM) by targeting a threonine residue. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	$ \begin{bmatrix} N \\ N \\ 0 \\ 0 \\ D \\ D$	bpV(phen), a insulin-mimetic agent, is a potent protein tyrosine phosphatase (PTP) and PTEN inhibitor with IC_{so} s of 38 nM, 343 nM and 920 nM for PTEN, PTP- β and PTP-1B, respectively. bpV(phen) inhibits proliferation of the protozoan parasite Leishmania in vitro. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	О 0 0 0 0 0 0 0 0 0 К ⁺
bpV(phen) trihydrate	Cat. No.: HY-122818	BR102375	Cat. No.: HY-128344
bpV(phen) trihydrate, a insulin-mimetic agent, is a potent protein tyrosine phosphatase (PTP) and PTEN inhibitor with IC ₅₀ s of 38 nM, 343 nM and 920 nM for PTEN, PTP- β and PTP-1B, respectively. Purity: \geq 98.0% Clinical Data: No Development Reported	N N H20 H20 0°,0°. К ⁺ H20 H20 H20	BR102375 is a non-TZD peroxisome proliferator-activated receptor γ (PPAR γ) full agonist for the treatment of type 2 diabetes, reveals EC ₅₀ value of 0.28 μ M and A _{max} ratio of 98%.Purity:>98%Clinical Data:No Development Reported	
Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg		Size: 1 mg, 5 mg	



Brusatol		BS-181	
(NSC 172924)	Cat. No.: HY-19543		Cat. No.: HY-13266
Brusatol (NSC 172924) is a unique inhibitor of the Nrf2 pathway that sensitizes a broad spectrum of cancer cells to Cisplatin and other chemotherapeutic agents. Brusatol enhances the efficacy of chemotherapy by inhibiting the Nrf2-mediated defense mechanism.Purity:99.89%Clinical Data:No Development Reported Size:10 mM × 1 mL, 5 mg, 10 mg		BS-181 is a potent and selective CDK7 inhibitor $(IC_{s_0}=21 \text{ nM})$ than Seliciclib (HY-30237). BS-181 is also against CDK2, CDK5 and CDK9 with IC_{s_0} values of 880, 3000 and 4200 nM, respectively (fails to block CDK1, 4 and 6).Purity:98.10%Clinical Data:No Development Reported Size:10 mM × 1 mL, 5 mg, 10 mg, 50 mg	HN HN HN HN
BTdCPU		BTK-IN-7	
	Cat. No.: HY-118266		Cat. No.: HY-143900
BTdCPU is a potent heme-regulated eIF2α kinase (HRI) activator. BTdCPU promotes eIF2α phosphorylation and induced apoptosis in resistant cell.		BTK-IN-7 is a potent and selective inhibitor of BTK (IC ₅₀ =4.0 nM). BTK-IN-7 has high selectivity in both enzymatic (ITK >250-fold, EGFR >2500-fold) and cellular levels(ITK >227-fold, EGFR 27-fold). BTK-IN-7 also has potent antitumor activity.	Children Chi
Purity: 99.48% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	10 mg	Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	
BTK-IN-9	Cat. No. 11/ 11/044	BIR-1	C-+ N- + UV 111(17
BTK-IN-9 is a reversible BTK inhibitors with potent antiproliferative activity in mantle cell lymphoma. BTK-IN-9 specifically disturbs mitochondrial membrane potential and increases reactive oxygen species level in Z138 cells. Purity: >98% Clinical Data: No Development Reported	$(\mathbf{x}_{1},\mathbf{N}_{0},\mathbf{N}_{1},$	BTR-1 is an active anti-cancer agent, causes S phase arrest, and affects DNA replication in leukemic cells. BTR-1 activates apoptosis and induces cell death. Purity: 99.96% Clinical Data: No Development Reported	
Size: 1 mg, 5 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	00 mg
BTSA1	C-4 No - UV 122054	BTZO-1	C-4 No - UV 110024
BTSA1 is a potent, high affinity and orally activeBAX activator with an IC_{so} of 250 nM and an EC_{so} of 144 nM. BTSA1 binds with high affinityand specificity to the N-terminal activation siteand induces conformational changes to BAX leadingto BAX-mediated apoptosis.Purity:99.74%		BTZO-1 binds to Macrophage migration inhibitory factor (MIF) with a K _a value of 68.6 nM, and its binding requires the N-terminal Pro1. Purity: 99.57%	
Clinical Data:No Development ReportedSize:5 mg, 10 mg, 50 mg, 100 mg		Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg	
Bucladesine calcium (Dibutyryl cAMP calcium salt; DBcAMP calcium salt)	Cat. No.: HY-B0764A	Bufarenogin	Cat. No. : HY-N6573
Bucladesine calcium salt (Dibutyryl-cAMP calcium salt;DC2797 calcium salt) is a cell-permeable cyclic AMP (cAMP) analog and selectively activates cAMP dependent protein kinase (PKA) by increasing the intracellular level of cAMP.		Bufarenogin induces intrinsic apoptosis via Bax and ANT cooperation.	HO TH HOH
Purity: 95.73% Clinical Data: Launched Size: 10 mM × 1 mL, 50 mg, 100 mg		Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg	



c-Met/HDAC-IN-2		c-Met/MEK1/Flt-3-IN-1	
	Cat. No.: HY-143462		Cat. No.: HY-145865
c-Met/HDAC-IN-2 is a highly potent c-Met and HDAC dual inhibitor with IC ₅₀ s of 18.49 nM and 5.40 nM for HDAC1 and c-Met, respectively. c-Met/HDAC-IN-2 has antiproliferative activities against certain cancer cell lines. Purity: >98% Clinical Data: No Development Reported	ەرىپىرىنى مىرچىن	Antiproliferative against-3 (comp 33) shows a prominent activity against Hela ($IC_{50} = 0.21 \mu M$), A549 ($IC_{50} = 0.39 \mu M$), and MCF-7 ($IC_{50} = 0.33 \mu M$), respectively. Antiproliferative against-3 (comp 33) also dose dependently induces apoptosis by arresting A549 cells at G1 phase. Purity: >98% Clinical Data: No Development Reported	.0-2000 ²²⁶ 0,
Size: 1 mg, 5 mg		Size: 1 mg, 5 mg	
c100.0			
C188-9	Cat No + HV 112209	C2 Ceramide	Cat No : HV 101190
(111-101)	Cat. No.: HY-112288	(Ceramide 2)	Cat. No.: HY-101180
C188-9 (TTI-101) is a STAT3 inhibitor, with a K _d of 4.7 nM. C188-9 inhibits G-CSF-induced STAT3 activation and STAT3-dependent gene expression. C188-9 induces apoptosis in AML cell lines and primary samples and inhibits colony formation by primary AML blasts.	о о о о н	C2 Ceramide (Ceramide 2) is the main lipid of the stratum corneum and a protein phosphatase 1 (PP1) activator. C2 Ceramide activates PP2A and ceramide-activated protein phosphatase (CAPP) .	No John
Purity: 99.90%	NO -	Purity: ≥98.0%	
Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	00 mg	Size: 5 mg	
C646		C8-Ceramide	
	Cat. No.: HY-13823	(N-Octanoyl-D-erythro-sphingosine)	Cat. No.: HY-108391
C646 is a selective and competitive histone acetyltransferase p300 inhibitor with K_i of 400 nM, and is less potent for other acetyltransferases.	ЛЦ ^й ю СДР ^о рон	C8-Ceramide (N-Octanoyl-D-erythro-sphingosine) is a cell-permeable analog of naturally occurring ceramides. C8-Ceramide has anti-proliferation properties and acts as a potent chemotherapeutic agent.	~~
Purity: ≥98.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 50 mg	0	Purity: ≥98.0% Clinical Data: No Development Reported Size: 5 mg	
CA 50		64224	
CA-5f	Cat. No. 11V 112609	CA224	Cot No. 11V 111207
	Cat. No.: HY-112698		Cat. No.: HY-111207
CA-5f is a potent late-stage macroautophagy/autophagy inhibitor via inhibiting autophagosome-lysosome fusion. CA-5f increases LC3B-II (a marker to monitor autophagy) and SQSTM1 protein, and also increases ROS production. Anti-tumor activity.	AND N CLO	CA224 (Compound 1) is a selective and orally active Cdk4–cyclin D1 inhibitor with an IC _{so} of 6.2 μM. CA224 induces cell apoptosis and shows antitumor activity.	
Purity: 99.40%		Purity: >98%	
Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10)0 mg	Size: 1 mg, 5 mg	
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Cabozantinib		Cabozantinib S-malate	
(XL184; BMS-907351)	Cat. No.: HY-13016	(XL184 S-malate; BMS-907351 S-malate)	Cat. No.: HY-12044
Cabozantinib is a potent multiple receptor tyrosine kinases (RTKs) inhibitor that inhibits VEGFR2, c-Met, Kit, Axl and FIt3 with IC ₅₀ s of 0.035, 1.3, 4.6, 7 and 11.3 nM, respectively.		Cabozantinib S-malate (XL184 S-malate) is a potent multiple receptor tyrosine kinases inhibitor that inhibits VEGFR2, c-Met, Kit, Axl and Flt3 with IC_{so} of 0.035, 1.3, 4.6, 7 and 11.3 nM, respectively.	
Purity: 99.96%		Purity: 99.84%	
Clinical Data: Launched Size: 10 mM × 1 mL, 5 mg, 10 mg. 50 mg. 100 mg. 2	200 mg	Clinical Data: Launched Size: 10 mM × 1 mL, 5 mq, 10 ma. 50 ma. 100 ma.	200 mg
, o		, J	

Coherentinih d4		Coherentinih dG	
Cabozantinid-04 (XI 184-d4: BMS-907351-d4)	Cat. No : HY-1301651	Cabozantinio-do	Cat No : HY-130165
Cabozantinib-d4 is deuterium labeled Cabozantinib. Cabozantinib is a potent multiple receptor tyrosine kinases (RTKs) inhibitor that inhibits VEGFR2, c-Met, Kit, Axl and Flt3 with IC50s of 0.035, 1.3, 4.6, 7 and 11.3 nM, respectively. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Cabozantinib-d6 (XL184-d6) is the deuterium labeled Cabozantinib. Cabozantinib is a potent multiple receptor tyrosine kinases (RTKs) inhibitor that inhibits VEGFR2, c-Met, Kit, Axl and FH3 with IC $_{so}$ s of 0.035, 1.3, 4.6, 7 and 11.3 nM, respectively.Purity:98.14% Clinical Data:No Development Reported Size:5 mg, 10 mg	
Cadein1	Cat. No. : HY-131143	Caffeic acid phenethyl ester	Cat. No.: HY-N0274
Cadein1, an isoquinolinium derivative, leads to a G2/M delay and caspase-dependent apoptosis in cancer cells with non- functional p53.		Caffeic acid phenethyl ester is a NF-κB inhibitor.	норосто
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:98.19%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 100 mg	
Calcimycin (A-23187; Antibiotic A-23187)	Cat. No.: HY-N6687	Calcimycin hemicalcium salt (A-23187 hemicalcium s Antibiotic A-23187 hemicalcium salt)	alt; Cat. No.: HY-N6687A
Calcimycin (A-23187) is an antibiotic and a unique divalent cation ionophore (like calcium and magnesium). Calcimycin induces Ca ²⁺ -dependent cell death by increasing intracellular calcium concentration. Calcimycin inhibits the growth of Gram-positive bacteria and some fungi. Purity: 99.56% Clinical Data: Phase 3 Size: 10 mM × 1 mL, 1 mg, 5 mg	HOC - CO-	Calcimycin hemicalcium salt (A-23187 hemicalcium salt) is an antibiotic and a unique divalent cation ionophore (like calcium and magnesium). Calcimycin hemicalcium salt induces Ca ²⁺ -dependent cell death by increasing intracellular calcium concentration. Purity: >98% Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg	HACCH HACCH
Calcimycin hemimagnesium		Calcium dobesilate	
(A-23187 hemimagnesium; Antibiotic A-23187 hemimagne	esiu@a)t. No.: HY-N6687B		Cat. No.: HY-111603
Calcimycin (A-23187) hemimagnesium is an antibiotic and a unique divalent cation ionophore (like calcium and magnesium). Calcimycin hemimagnesium induces Ca ²⁺ -dependent cell death by increasing intracellular calcium concentration. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	4,2000,000,000 h-1,000,000,000 h-1,000,000,000	Calcium dobesilate, a vasoprotective, is widely used in chronic venous disease, diabetic retinopathy and the symptoms of haemorrhoidal attack in many countries. Purity: ≥98.0% Clinical Data: Launched Size: 10 mM × 1 mL, 500 mg	$HO = \begin{pmatrix} 0, 0 \\ 0 \\ 0 \\ 0 \\ HO \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ $
Calicheamicin		CALP1	
Calicheamicin, an antitumor antibiotic, is a cytotoxic agent that causes double-strand DNA breaks. Calicheamicin is a DNA synthesis inhibitor. Purity: 98.28% Clinical Data: No Development Reported Size: 1 mg, 5 mg	Cat. No.: HY-19609	CALP1 is a calmodulin (CaM) agonist (K_d of 88 μ M) with binding to the CaM EF-hand/Ca ²⁺ -binding site. CALP1 blocks calcium influx and apoptosis (IC_{so} of 44.78 μ M) through inhibition of calcium channel opening.Purity:>98% Clinical Data:No Development Reported Size:1 mg, 5 mg	Cat. No.: HY-P10/7

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		Colportin	
CALCI IFA	Cat. No.: HY-P1077A	Caipeptin	Cat. No. : HY-100223
CALP1 TFA is a calmodulin (CaM) agonist (K_d of 88 μ M) with binding to the CaM EF-hand/Ca ²⁺ -binding site. CALP1 TFA blocks calcium influx and apoptosis (IC_{so} of 44.78 μ M) through inhibition of calcium channel opening.	suggeder.	Calpeptin is a potent, cell penetrating calpain inhibitor, with an ID_{s0} of 40 nM for Calpain I in human platelets. Calpeptin is also an inhibitor of cathepsin K .	Coluctor
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity: ≥98.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	00 mg
Calphostin C (UCN-1028C)	Cat. No.: HY-105416	Calycosin (Cyclosin)	Cat. No.: HY-N0519
Calphostin C is a potent and specific inhibitor of protein kinase C . Calphostin C is an antitumor antibiotic. Calphostin C has 1000 times more inhibitory to protein kinase C with an IC_{so} of 0.05 μ M than other protein kinases.		Calycosin (Cyclosin) is a natural active compound with anti-oxidative and anti-inflammation activity. IC50 value: Target: in vitro: calycosin had obvious anti-proliferation effects on SKOV3 cells in a dose- and time-dependent manner.	HO CONTRACTOR
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	но~~ор о~	Purity: 99.89% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg	
Cambinol	Cat. No.: HY-100732	Camellianin A	Cat. No.: HY-N2298
Cambinol is a SIRT1 and SIRT2 inhibitor with IC_{s0} values of 56 μ M and 59 μ M, respectively. Cambinol is a potent brain penetrant neutral sphingomyelinase (N-SMase) inhibitor (exosome inhibitor).	NH NH	Camellianin A, the main flavonoid in A. nitida leaves, displays anticancer activity and angiotensin converting enzyme (ACE)-inhibitory activity.	$H_{0} \stackrel{1}{\overset{1}{\overset{1}{\overset{1}{\overset{1}{\overset{1}{\overset{1}{\overset{1}{$
Purity: 99.70% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	σ
Camptothecin (Campathecin; (S)-(+)-Camptothecin; CPT)	Cat. No.: HY-16560	Camptothecin-d5 (Campathecin-d5; (S)-(+)-Camptothecin-d5; CPT-d5)	Cat. No. : HY-16560S
Camptothecin (CPT), a kind of alkaloid, is a DNA topoisomerase I (Topo I) inhibitor with an IC _{so} of 679 nM.		Camptothecin-d5 (Campathecin-d5) is the deuterium labeled Camptothecin. Camptothecin (CPT), a kind of alkaloid, is a DNA topoisomerase I (Topo I) inhibitor with an IC ₅₀ of 679 nM.	CTN DHOLO O
Purity: 99.69% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 500 mg	. 10 0	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	U U
Candesartan Cilexetil (TCV-116)	Cat. No.: HY-17505	Cantrixil (TRX-E-002-1)	Cat. No. : HY-114250
Candesartan Cilexetil (TCV-116) is an angiotensin II receptor antagonist used mainly for the treatment of hypertension.	J ^N o ^N N ^N N ^N O ^N O ^N O ^N O ^N O ^N O	Cantrixil (TRX-E-002-1), an active enantiomer of TRX-E-002, is a second-generation super-benzopyran (SBP) compound. Cantrixil increases phosphorylated c-Jun levels resulting in caspase-mediated apoptosis in ovarian cancer cells.	HO LO COL
Purity: 99.77% Clinical Data: Launched Size: 10 mM × 1 mL, 500 mg, 1 g	Q	Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	он

Capecitabine		Capecitabine-d11	
Capecitabine is an oral prodrug that is converted to its active metabolite, 5-FU, by thymidine phosphorylase.		Capecitabine-d11 is the deuterium labeled Capecitabine. Capecitabine is an oral prodrug that is converted to its active metabolite, 5-FU, by thymidine phosphorylase.	
Purity:99.73%Clinical Data:LaunchedSize:10 mM × 1 mL, 500 mg, 1 g, 5 g		Purity:>98%Clinical Data:Size:1 mg, 5 mg	
Capmatinib (INC280; INCB28060) Ca	at. No.: HY-13404	Capsaicin ((E)-Capsaicin)	Cat. No.: HY-10448
Capmatinib (INC280; INCB28060) is a potent, orally active, selective, and ATP competitive c-Met kinase inhibitor (IC ₅₀ =0.13 nM).		Capsaicin ((E)-Capsaicin), an active component of chili peppers, is a TRPV1 agonist. Capsaicin has pain relief, antioxidant, anti-inflammatory, neuroprotection and anti-cancer effects.	-ogget g
Purity: 99.92% Clinical Data: Launched Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg/t	g	Purity: 99.85% Clinical Data: Launched Size: 10 mM × 1 mL, 50 mg, 100 mg	
Capsaicin-d3 ((E)-Capsaicin-d3) Cat.	No.: HY-10448S1	Capsaicinoid	Cat. No.: HY-10448A
Capsaicin-d3 ((E)-Capsaicin-d3) is the deuterium labeled Capsaicin. Capsaicin ((E)-Capsaicin), an active component of chili peppers, is a TRPV1 agonist. Capsaicin has pain relief, antioxidant, anti-inflammatory, neuroprotection and anti-cancer effects.	g#~CCote	Capsaicinoid is a mixture of Capsaicin and Dihydrocapsaicin. Capsaicinoid is an capsaicin receptor (TRPV1) agonist.	ogget times
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 10 mg		Purity:99.46%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 50 mg	
Capsazepine	at. No. : HY-15640	Cardanol monoene (Cardanol C15:1)	Cat. No. : HY-119979
Capsazepine is a synthetic analogue of the sensory neurone excitotoxin, and an antagonist of TRPV1 receptor with an IC ₅₀ of 562 nM.	Colt-O°	Cardanol monoene (Cardanol C15:1) is a phenolic compound which can be found in cashew nut shell liquid. Cardanol monoene can induce mitochondria-associated apoptosis in human melanoma cells.	~~~~Q.,
Purity:99.17%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg, 50 mg, 100 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
Carfilzomib (PR-171) Co	at. No. : HY-10455	Carfilzomib-d8	Cat. No. : HY-10455S
Carfilzomib (PR-171) is an irreversible proteasome inhibitor with an IC_{so} of 5 nM in ANBL-6 and RPMI 8226 cells.	hinite.	Carfilzomib-d8 is deuterium labeled Carfilzomib. Carfilzomib (PR-171) is an irreversible proteasome inhibitor with an IC50 of 5 nM in ANBL-6 and RPMI 8226 cells.	
Purity: 99.96% Clinical Data: Launched Size: 5 mg, 10 mg, 50 mg, 100 mg, 200 mg	199 8 /201	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	

Carnosic acid	Cat No : HY-N0644	Carubicin (Carminomycin: Carminomicin I)	Cat No: HY-B2171
Carnosic acid has demonstrated inhibition of oxidative stress and inflammation, suppression of cell proliferation, and antibacterial activity. Purity: 99.15% Clinical Data: No Development Reported Size: 10 mg, 50 mg		Carubicin (Carminomycin) is a microbially-derived compound. Carubicin is an effective inhibitor of VHL-defective (VHL-/-) CCRCC cell proliferation. Carubicin also induces apoptosis by a mechanism independent of p53 or hypoxia-inducible factor HIF2. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	OH O OH OF NH2
Carubicin hydrochloride (Carminomycin hydrochloride; Carminomicin I hydrochlorid	e) Cat. No.: HY-B2171A	Carvacrol	Cat. No.: HY-N0711
Carubicin hydrochloride is a microbially-derived compound. Carubicin hydrochloride is an effective inhibitor of VHL-defective (VHL-/-) CCRCC cell proliferation. Carubicin hydrochloride also induces apoptosis by a mechanism independent of p53 or hypoxia-inducible factor HIF2. Purity: 98.67% Clinical Data: No Development Reported Size: 5 mg, 10 mg, 50 mg		Carvacrol is a monoterpenoid phenol isolated from Lamiaceae family plants, with antioxidant, anti-inflammatory and anticancer properties. Carvacrol causes cell cycle arrest in G0/G1, downregulates Notch-1, and Jagged-1, and induces apoptosis. Purity: 99.96% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 100 mg	но
Casein Kinase inhibitor A51	Cat. No.: HY-123954	Casein Kinase inhibitor A86	Cat. No.: HY-123955
Casein Kinase inhibitor A51 is a potent and orally active casein kinase 1α (CK1α) inhibitor. Casein Kinase inhibitor A51 induces leukemia cell apoptosis, and has potent anti-leukemic activities. Purity: 98.42% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10		Casein Kinase inhibitor A86 is a potent and orally active casein kinase 1α (CK1α) inhibitor. Casein Kinase inhibitor A86 also inhibits of CDK7 (TFIIH) and CDK9 (P-TEFb). Casein Kinase inhibitor A861 induces leukemia cell apoptosis, and has potent anti-leukemic activities.Purity:98.47% Clinical Data:No Development Reported Size:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
Catachin		Condetin	
((+)-Catechin; Cianidanol; Catechuic acid)	Cat. No.: HY-N0898	Caudatin	Cat. No.: HY-N1983
Catechin ((+)-Catechin) inhibits cyclooxygenase-1 (COX-1) with an IC ₅₀ of 1.4 μ M.	но строн он	Caudatin is a steroidal cmpound found in Cynanchum auriculatum, causes cell cycle arrest and induces apoptosis , with anti-cancer and antiangiogenic properties.	оториски странации и с
Clinical Data: Launched Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg		Clinical Data: No Development Reported Size: 5 mg, 10 mg	
CAY10404	Cat. No.: HY-121537	CAY10505	Cat. No.: HY-13530
CAY10404 is a potent and selective cyclooxygenase-2 (COX-2) inhibitor with an IC_{s0} of 1 nM and a selectivity index (SI; COX-1 IC_{s0} /COX-2 IC_{s0}) of > 500000.	N. C. C.	CAY10505 is a potent and selective $PI3K\gamma$ inhibitor with an IC_{s0} of 30 nM in neurons.	F.C. C. S.
Purity:99.79%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	F K 5	Purity:99.75%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg, 25 mg, 50 mg, 100 mg	



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CDK0/PIM1-IN-1	Cat. No.: HY-142696	CDK9-IN-7	Cat. No.: HY-126251
$\begin{array}{llllllllllllllllllllllllllllllllllll$	"O CLASS	CDK9-IN-7 (compound 21e) is a selective, highly potent, and orally active CDK9/cyclin T inhibitor (IC _{so} =11 nM), which exhibits more potent over other CDKs (CDK4/cyclinD=148 nM; CDK6/cyclinD=145 nM). CDK9-IN-7 shows antitumor activity without obvious toxicity. Purity: 99.81% Clinical Data: No Development Reported Size: 5 mg, 10 mg, 50 mg	State of the second
CDKI-73 (LS-007)	Cat. No.: HY-12445	Cearoin	Cat. No.: HY-N8418
CDKI-73 (LS-007) is an orally active and highly efficacious CDK9 inhibitor, with K _i values of 4 nM, 4 nM and 3 nM for CDK9, CDK1 and CDK2, respectively. CDKI-73 down-regulates the RNAPII phosphorylation.	HN S NN HANGO	Cearoin increases autophagy and apoptosis through the production of ROS and the activation of ERK .	но
Purity:99.58%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity: ≥98.0% Clinical Data: No Development Reported Size: 1 mg	94050 (CONEQA
Caramida C6 d7		Coronib 2	
Ceramide Co-d7	Cat. No.: HY-19542S	Ceramo-2	Cat. No.: HY-116147
Ceramide C6-d7 is the deuterium labeled Ceramide C6. Ceramide C6	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Ceranib-2 is a potent and nonlipid ceramidase inhibitor that inhibits cellular ceramidase activity with an IC_{s0} of 28 μ M in SKOV3 cells.	
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Purity:99.25%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	~
CFM-4		CGP 57380	
	Cat. No.: HY-103255		Cat. No.: HY-10520
CFM-4 is a potent small molecular antagonist of CARP-1/APC-2 binding. CFM-4 prevents CARP-1 binding with APC-2, causes G ₂ M cell cycle arrest, and induces apoptosis with an IC ₅₀ range of 10-15 µM. CFM-4 also suppresses growth of drug-resistant human breast cancer cells. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		CGP 57380 is a cell-permeable pyrazolo-pyrimidine compound that acts as a selective inhibitor of Mnk1 with ICs0 of 2.2 μ M, but has no inhibitory activity against p38, JNK1, ERK1/2, PKC, or Src-like kinases.Purity:98.0% Clinical Data:No Development Reported Size:10 mM × 1 mL, 10 mg, 50 mg	
Chaetoglobosin A	Cat. No.: HY-N6744	Chelerythrine	Cat. No.: HY-N2359
Chaetoglobosin A, the active principle within the extract of Penicillium aquamarinium, is a member of the cytochalasan family. Chaetoglobosin A preferentially induces apoptosis .		Chelerythrine is a natural alkaloid, acts as a potent and selective Ca ²⁺ /phospholopid-dependent PKC antagonist, with an IC ₅₀ of 0.7 μ M. Chelerythrine has antitumor, antidiabetic and anti-inflammatory activity.	STOR STOR
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 20 mg	







CI-amidine TFA	Cat. No.: HY-100574B	Cladribine (2-Chloro-2'-deoxyadenosine; CldAdo; 2CdA)	Cat. No.: HY-13599
$ \begin{array}{llllllllllllllllllllllllllllllllllll$		Cladribine (2-Chloro-2'-deoxyadenosine), a purine nucleoside analog, is an orally active adenosine deaminase inhibitor. Cladribine functions as an inhibitor of DNA synthesis to block the repair of the damaged DNA. Cladribine can inhibit DNA methylation. Purity: 99.97% Clinical Data: Launched Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg	
CLEFMA	Cat. No. : HY-136718	Clitocine	Cat. No.: HY-118341
CLEFMA is a curcuminoid with antitumor activity. CLEFMA inhibits tumor growth is associated with NF-κB-regulated anti-inflammatory and anti-metastatic effects. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Clitocine, an adenosine nucleoside analog isolated from mushroom, is a potent and efficacious readthrough agent. Clitocine acts as a suppressor of nonsense mutations and can induce the production of p53 protein in cells harboring p53 nonsense-mutated alleles. Purity: 95.88% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	но но он о
Clobenpropit dihydrobromide	Cat. No. : HY-101198	Clofarabine	Cat. No. : HY-A0005
Clobenpropit dihydrobromide is a potent histamine H3R antagonist/inverse agonist with a pEC ₅₀ of 8.07 for histamine H3LR. Clobenpropit dihydrobromide acts as partial agonist at histamine H4 receptors (K ₁ 13 nM).		Clofarabine, a nucleoside analogue for research of cancer, is a potent inhibitor of ribonucleotide reductase (IC_{50} =65 nM) by binding to the allosteric site on the regulatory subunit.	
Purity: >98% Clinical Data: No Development Reported Size: 5 mg		Purity: 99.09% Clinical Data: Launched Size: 10 mM × 1 mL, 10 mg, 50 mg	но
Clofilium tosylate	Cat. No. : HY-33350	Clovamide (trans-Clovamide)	Cat. No.: HY-122267
Clofilium tosylate, a potassium channel blocker, induces apoptosis of human promyelocytic leukemia (HL-60) cells via Bcl-2-insensitive activation of caspase-3. Antiarrhythmic agent.	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Clovamide (trans-Clovamide), a natural phenolic compound, is a potent antioxidant. Clovamide is an excellent ROS and oxygen radical scavenger. Clovamide also has anti-inflammatory and neuroprotective effects.	но страни страни
Purity: \geq 98.0%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg, 50 mg		Purity:98.48%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
CM-272	Cat. No. : HY-101925	CMC2.24 (TRB-N0224)	Cat. No.: HY-120793
CM-272 is a first-in-class, potent, selective, substrate-competitive and reversible dual G9a/DNA methyltransferases (DNMTs) inhibitor with antitumor activities.	another the	CMC2.24 (TRB-N0224), an orally active tricarbonylmethane agent, is effective against pancreatic tumor in mice by inhibiting Ras activation and its downstream effector ERK1/2 pathway.	HO, C, J, H, C, OH
Purity: 99.27% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity: 96.48% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	00 mg

CMID 2		CMI D010509	
CMLD-2	Cat. No.: HY-124828	(SDS-1-021)	Cat. No.: HY-119271
CMLD-2, an inhibitor of HuR-ARE interaction , competitively binds HuR protein disrupting its interaction with adenine-uridine rich elements (ARE)-containing mRNAs (K _i =350 nM).		CMLD010509 (SDS-1-021) is a highly specific inhibitor of the oncogenic translation program supporting multiple myeloma (MM)-including key oncoproteins such as MYC, MDM2, CCND1, MAF, and MCL-1.	
Clinical Data: No Development Reported Size: 5 mg, 10 mg		Clinical Data: No Development Reported Size: 1 mg, 5 mg	
Cobimetinib (GDC-0973; XL518)	Cat. No. : HY-13064	Cobimetinib hemifumarate (GDC-0973 hemifumarate; XL-518 hemifumarate)	Cat. No.: HY-13064A
Cobimetinib (GDC-0973, RG7420) is a potent, selective and oral MEK1 inhibitor with an IC_{so} of 4.2 nM for MEK1.		Cobimetinib hemifumarate is a novel selective MEK1 inhibitor, and the IC ₅₀ value against MEK1 is 4.2 nM.	
Purity: 99.71% Clinical Data: Launched Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	Ť	Purity: 98.08% Clinical Data: Launched Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg	Ť
Cobimetinib-d4 (GDC-0973-d4; XL518-d4)	Cat. No. : HY-13064S	Coenzyme Q9 (Ubiquinone Q9; CoQ9; Ubiquinone 9)	Cat. No .: HY-101415
Cobimetinib-d4 (GDC-0973-d4) is the deuterium labeled Cobimetinib. Cobimetinib (GDC-0973, RG7420) is a potent, selective and oral MEK1 inhibitor with an IC_{50} of 4.2 nM for MEK1 .		Coenzyme Q9 (Ubiquinone Q9), the major form of ubiquinone in rodents, is an amphipathic molecular component of the electron transport chain that functions as an endogenous antioxidant. Coenzyme Q9 attenuates the diabetes-induced decreases in antioxidant defense mechanisms.	
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	Y	Purity:≥98.0%Clinical Data:No Development ReportedSize:1 mg, 5 mg, 10 mg	
COG1410		Colchicine	
	Cat. No.: HY-P2136		Cat. No.: HY-16569
COG1410 is an apolipoprotein E-derived peptide. COG1410 exerts neuroprotective and antiinflammatory effects in a murine model of traumatic brain injury (TBI). COG1410 can be used for the research of neurological disease.	Ac-AS-(Aib)-LRKL-(Aib)-KRLL-NH2	Colchicine is a tubulin inhibitor and a microtubule disrupting agent. Colchicine inhibits microtubule polymerization with an IC ₅₀ of 3 nM. Colchicine is also a competitive antagonist of the a3 glycine receptors (GlyRs).	
Purity:99.49%Clinical Data:No Development ReportedSize:1 mg, 5 mg, 10 mg		Purity: 99.87% Clinical Data: Launched Size: 10 mM × 1 mL, 200 mg, 500 mg	у-NH O
Colchicine-d3	Cat. No. : HY-16569S1	Colchicine-d6	Cat. No.: HY-16569S
$\begin{array}{llllllllllllllllllllllllllllllllllll$		$ Colchicine-d6 is the deuterium labeled Colchicine. Colchicine is a tubulin inhibitor and a microtubule disrupting agent. Colchicine inhibits microtubule polymerization with an IC50 of 3 nM. Colchicine is also a competitive antagonist of the \alpha3 glycine receptors (GlyRs). Purity: >98% Clinical Data: No Development Reported$	
Size: 1 mg, 5 mg, 10 mg		Size: 1 mg, 10 mg	

Colivelin TFA		Columbianadin	
	Cat. No.: HY-P1061A		Cat. No.: HY-N0362
Colivelin TFA is a brain penetrant neuroprotective peptide and a potent activator of STAT3 , suppresses neuronal death by activating STAT3 in vitro.	SALLRSPAPAGASRILLLTGEDUP (TFA safe	Columbianadin, a natural coumarin from, is known to have various biological activities including anti-inflammatory and anti-cancer effects.	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
Purity:99.22%Clinical Data:No Development ReportedSize:500 μg, 1 mg		Purity:99.03%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg	
Concanavalin A	Cat. No.: HY-P2149	Conglobatin (FW-04-806)	Cat. No. : HY-119906
Concanavalin A is a Ca ²⁺ /Mn ²⁺ -dependent and mannose/glucose-binding plant lectin that can be found in jack bean. Concanavalin A can induce programmed cell death.	Concanavalin A	Conglobatin (FW-04-806), a macrolide dilactone, is isolated from the culture of Streptomyces conglobatus. Conglobatin is an orally active Hsp90 inhibitor. Conglobatin can bind to the N-terminal domain of Hsp90 and disrupt Hsp90-Cdc37 complex formation.	NECHICAN Services
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity: >98% Clinical Data: No Development Reported Size: 500 μg, 1 mg, 5 mg	
Coniferaldehyde (Ferulaldehyde)	Cat. No.: HY-N2535	Conophylline	Cat. No.: HY-N3619
Coniferaldehyde (Ferulaldehyde) is an effective inducer of heme oxygenase-1 (HO-1). Coniferaldehyde exerts anti-inflammatory properties in response to LPS.	HO	Conophylline is a vinca alkaloid extracted from leaves of a tropical plant Ervatamia microphylla. Conophylline is a differentiation inducer of for pancreatic cells. Conophylline suppresses HSC and induces apoptosis.	A CONTRACTOR
Purity:99.94%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity: ≥98.0% Clinical Data: No Development Reported Size: 1 mg, 5 mg	0
Conanlisih		Copaniisib dibydrochloride	
(BAY 80-6946)	Cat. No.: HY-15346	(BAY 80-6946 dihydrochloride)	Cat. No.: HY-15346A
Copanlisib (BAY 80-6946) is a potent, selective and ATP-competitive pan-class I PI3K inhibitor, with IC ₅₀ s of 0.5 nM, 0.7 nM, 3.7 nM and 6.4 nM for PI3K α , PI3K β , PI3K β and PI3K γ , respectively.	Cu-change Canada	Copanlisib dihydrochloride (BAY 80-6946 dihydrochloride) is a potent, selective and ATP-competitive pan-class I PI3K inhibitor, with IC ₅₀ s of 0.5 nM, 0.7 nM, 3.7 nM and 6.4 nM for PI3K α , PI3K β , PI3K β and PI3K γ , respectively.	0 (
Purity: 99.50% Clinical Data: Launched Size: 5 mg, 10 mg, 50 mg, 100 mg		Purity: 99.55% Clinical Data: Launched Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	-0.4 SPN4
Copanlisib-d6 (BAY 80-6946-d6)	Cat. No.: HY-15346S1	Copanlisib-d8 (BAY 80-6946-d8)	Cat. No.: HY-15346S
Copanlisib-d6 (BAY 80-6946-d6) is the deuterium labeled Copanlisib. Copanlisib (BAY 80-6946) is a potent, selective and ATP-competitive pan-class I PI3K inhibitor, with $IC_{so}s$ of 0.5 nM, 0.7 nM, 3.7 nM and 6.4 nM for PI3K α , PI3K β and PI3K γ , respectively. Purity: >98%		Copanlisib-d8 (BAY 80-6946-d8) is the deuterium labeled Copanlisib. Copanlisib (BAY 80-6946) is a potent, selective and ATP-competitive pan-class I PI3K inhibitor, with IC ₅₀ s of 0.5 nM, 0.7 nM, 3.7 nM and 6.4 nM for PI3K α , PI3K β , PI3K β and PI3K γ , respectively. Purity: >98%	$(\begin{array}{c} \begin{array}{c} \begin{array}{c} \\ \\ \end{array} \\ \\ \end{array} \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ $
Clinical Data:No Development ReportedSize:1 mg, 5 mg		Clinical Data: No Development Reported Size: 1 mg, 5 mg	



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Crebanine		cRIPGBM	
	Cat. No.: HY-N2255		Cat. No.: HY-125466
Crebanine, an alkaloid from Stephania venosa, induces G1 arrest and apoptosis in human cancer cells. Crebanine exhibits anti-inflammatory activity via suppressing MAPKs and Akt signaling. Crebanine also possesses antiarrhythmic effect. Purity: 99.54% Clinical Data: No Development Reported Size: 5 mg 10 mg 20 mg		cRIPGBM, a proapoptotic derivative of RIPGBM, a cell type-selective inducer of apoptosis in GBM cancer stem cells (CSCs) by binding to receptor-interacting protein kinase 2 (RIPK2), with an EC ₅₀ of 68 nM in GBM-1 cells. Purity: >98% Clinical Data: No Development Reported Size:	
Size: 5 mg, 10 mg, 20 mg		Size: 1 mg, 5 mg	
Crolibulin (EPC2407)	Cat. No. : HY-13603	CS1	Cat. No.: HY-137005
Crolibulin (EPC2407) is a tubulin polymerization inhibitor, with potent apoptosis induction and cell growth inhibition. Crolibulin has anti-tumor activity. Crolibulin also has cardiovascular toxicity and neurotoxicity.		CS1 is a potent DNA Topo II α inhibitor. CS1 displays broad-spectrum in vitro antitumor effects, low toxicity in vivo and potential anti-multidrug resistance capabilities. CS1 leads to DNA damage, cell cycle arrest at G2/M phase and apoptosis .	но
Purity: 98.99% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg	0~	Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	
CSRM617		СТВ	
	Cat. No.: HY-122611		Cat. No.: HY-134964
CSRM617 is a selective small-molecule inhibitor of the transcription factor ONECUT2 (OC2, a master regulator of androgen receptor) with a K _d of 7.43 uM in SPR assays, binding to OC2-HOX domain directly. CSRM617 induces apoptosis by appearance of cleaved Caspase-3 and PARP.	HO NH2 N NO OH	CTB is a potent p300 histone acetyltransferase activator. CTB can effectively induce apoptosis in MCF-7 cells.	
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Purity:99.76%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
		Cucurbitacia P	
0-5	Cat. No.: HY-121638		Cat. No.: HY-N0416
CU-3 is the racemate of (5Z,2E)-CU-3. (5Z,2E)-CU-3 is a potent and selective inhibitor against the α -isozyme of DGK with an IC _{so} value of 0.6 μ M, competitively inhibits the affinity of DGK α for ATP with a K _m value of 0.48 mM.	O HAL-N S	Cucurbitacin B belongs to a class of highly oxidized tetracyclic triterpenoids; could repress cancer cell progression.	
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg	13523	Purity: 99.92% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg	- Sear
Cucurbitacin IIa (Hemslecin A)	Cat. No.: HY-N1988	Cucurbitacin IIb	Cat. No.: HY-N1987
Cucurbitacin IIa is a triterpene isolated from Hemsleya amalils Diels, induces apoptosis of cancer cells, reduces expression of survivin , reduces phospho-Histone H3 and increases cleaved PARP in cancer cells.	HO HO L	Cucurbitacin IIb is an active component isolated from Hemsleya amabilis , induces apoptosis with anti-inflammatory activity.	HO HO HO HO
Purity:99.27%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 20 mg		Purity:98.87%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg	



Cysteamine hydrochloride (2-Aminoethanethiol hyd	rochloride;	Cysteamine-d4 hydrochloride (2-Aminoethanethiol	-d4
Cysteamine hydrochloride (2-Aminoethanethiol hydrochloride) is an orally active agent for the treatment of nephropathic cystinosis and an antioxidant. Purity: ≥95.0% Clinical Data: Launched Size: 10 mM × 1 mL, 500 mg, 5 g	HS NH ₂ HCI	Cysteamine-d4 (2-Aminoethanethiol-d4 hydrochloride) is the deuterium labeled Cysteamine hydrochloride. Cysteamine hydrochloride (2-Aminoethanethiol hydrochloride) is an orally active agent for the treatment of nephropathic cystinosis and an antioxidant. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	$\begin{array}{c} D & SH \\ D \longrightarrow D & HCI \\ H_2N & D \end{array}$
Cytarabine (Cytosine β-D-arabinofuranoside; Cytosine Arabinoside; Ara-C)	Cat. No .: HY-13605	Cytarabine-d2	Cat. No.: HY-13605S
Cytarabine, a nucleoside analog, causes S phase cell cycle arrest and inhibits DNA polymerase. Cytarabine inhibits DNA synthesis with an IC ₅₀ of 16 nM. Cytarabine has antiviral effects against HSV. Purity: 99.96% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 500 mg, 1 g		Cytarabine-d2 is the deuterium labeled Cytarabine. Cytarabine, a nucleoside analog, causes S phase cell cycle arrest and inhibits DNA polymerase. Cytarabine inhibits DNA synthesis with an IC ₅₀ of 16 nM. Cytarabine has antiviral effects against HSV. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	$ \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \end{array}\\ \end{array}\\ \end{array} \\ \begin{array}{c} \end{array}\\ \end{array} \\ \begin{array}{c} \end{array}\\ \end{array} \\ \begin{array}{c} \end{array} \\ \begin{array}{c} \end{array}\\ \end{array} \\ \begin{array}{c} \end{array} \\ \begin{array}{c} \end{array} \\ \end{array} \\ \begin{array}{c} \end{array} \\ \end{array} \\ \begin{array}{c} \end{array} \\ \end{array} \\ \end{array} \\ \begin{array}{c} \end{array} \\ \end{array} \\ \end{array} \\ \begin{array}{c} \end{array} \\ \end{array} \\ \end{array} \\ \end{array} \\ \end{array} \\ \begin{array}{c} \end{array} \\ \end{array} \\ \end{array} \\ \end{array} \\ \end{array} \\ \end{array} \\ \end{array} $ } \\ \end{array} \\ \end{array} \\ \end{array} } \\ \end{array} \\ \end{array} \\ \end{array} \\ \end{array} } \\ \end{array} \\ \end{array} \\ \end{array} } \\ \end{array} \\ \end{array} \\ \end{array} \\ \end{array} } \\ \end{array} } \\ \end{array} } \\ \end{array}
Cytostatin	Cat. No.: HY-113612	D-Cl-amidine	Cat. No.: HY-100574C
Cytostatin is a potent and selective inhibitor of PP₂A with promising antitumor activity. Cytostatin is also an inhibitor of cell adhesion to extracellular matrix and induces cell apoptosis . Cytostatin belongs to the fostriecin family of natural products.	Pri pristing	D-Cl-amidine is a potent and highly selective PAD1 inhibitor. D-Cl-amidine is well-torelated with no significant toxicity.	
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg		Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	
D-CI-amidine hydrochloride	Cat. No. : HY-100574D	D-Mannitol (Mannitol; Mannite)	Cat. No. : HY-N0378
D-Cl-amidine hydrochloride is a potent and highly selective PAD1 inhibitor. D-Cl-amidine is well-torelated with no significant toxicity.		D-Mannitol is an osmotic diuretic agent and a weak renal vasodilator. Target: Others D(-)Mannitol is a sugar alcohol that can be used as an inert osmotic control substance.	он он ноон он он он
Purity:99.40%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg		Purity: ≥98.0% Clinical Data: Launched Size: 10 mM × 1 mL, 500 mg, 5 g, 10 g	
D-Mannitol-d8 (Mannitol-d8; Mannite-d8)	Cat. No.: HY-N0378S	D-Pantothenic acid hemicalcium salt (Calcium par Calcium D-pantothenate; Vitamin B5 calcium salt)	ntothenate; Cat. No.: HY-N0681
D-Mannitol-d8 (Mannitol-d8) is the deuterium labeled D-Mannitol. D-Mannitol is an osmotic diuretic agent and a weak renal vasodilator.	HO DHO D D DHO	D-Pantothenic acid hemicalcium salt (Vitamin B5 calcium salt), a vitamin, can reduce the patulin content of the apple juice.	о Ц у Ц () Н он он 0.5Са ²⁺
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity: ≥95.0% Clinical Data: Launched Size: 10 mM × 1 mL, 500 mg, 5 g	

D-Pantothenic acid sodium		D4476	
(Sodium pantothenate; Vitamin B5 sodium)	Cat. No.: HY-B0430A	(Casein Kinase I Inhibitor)	Cat. No.: HY-10324
D-Pantothenic acid sodium (Sodium pantothenate) is an essential trace nutrient that functions as the obligate precursor of coenzyme A (CoA).		D4476 is a potent, selective and cell-permeable inhibitor of casein kinase 1(CK1) with an IC_{s0} value of 0.3 μM in vitro.	
Purity:≥98.0%Clinical Data:LaunchedSize:10 mM × 1 mL, 100 mg		Purity:99.51%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg	
Dabuzalgron		Dacarbazine	
(Ro 115-1240)	Cat. No.: HY-117071	(Imidazole Carboxamide)	Cat. No.: HY-B0078
Dabuzalgron (Ro 115-1240) is an orally active and selective α-1A adrenergic receptor agonist for the treatment of urinary incontinence. Dabuzalgron protects against Doxorubicin-induced cardiotoxicity by preserving mitochondrial function.Purity:98.72%Clinical Data:No Development Reported Size:10 mM × 1 mL, 5 mg, 10 mg	THE STREET STREE	Dacarbazine(DTIC-Dome; DTIC) is an antineoplastic agent. It has significant activity against melanomas. Purity: ≥98.0% Clinical Data: Launched Size: 10 mM × 1 mL, 200 mg, 1 g	
Dacarbazine-d6		Dacomitinib	
(Imidazole Carboxamide-d6)	Cat. No.: HY-B0078S	(PF-00299804; PF-299804)	Cat. No.: HY-13272
Dacarbazine-d6 (Imidazole Carboxamide-d6) is the deuterium labeled Dacarbazine. Dacarbazine(DTIC-Dome; DTIC) is an antineoplastic agent. It has significant activity against melanomas. Purity: >98% Clinical Data: No Development Reported		Dacomitinib (PF-00299804) is a specific and irreversible inhibitor of the ERBB family of kinases with IC ₅₀ s of 6 nM, 45.7 nM and 73.7 nM for EGFR, ERBB2, and ERBB4, respectively. Purity: 99.92% Clinical Data: Launched	
Size: 1 mg, 10 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg,	200 mg
Dacomitinib-d10 (PF-00299804-d10; PF-299804-d10)	Cat. No.: HY-13272S3	Dacomitinib-d10 dihydrochloride (PF-00299804-d dihydrochloride; PF-299804-d10 dihydrochloride)	10 Cat. No.: HY-13272S2
Dacomitinib-d10 is deuterium labeled Dacomitinib. Dacomitinib (PF-00299804) is a specific and irreversible inhibitor of the ERBB family of kinases with IC50s of 6 nM, 45.7 nM and 73.7 nM for EGFR, ERBB2, and ERBB4, respectively. Purity: >98% Clinical Data: No Development Reported Size: 1 mg.5 mg		Dacomitinib-d10 (PF-00299804-d10) dihydrochloride is the deuterium labeled Dacomitinib dihydrochloride. Purity: >98% Clinical Data: No Development Reported Size: 1 mg. 5 mg	HN C HO
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Dacomitinib-d3 (PF-00299804-d3; PF-299804-d3)	Cat. No.: HY-13272S	Dacomitinib-d5 (PF-00299804-d5; PF-299804-d5)	Cat. No.: HY-13272S1
Dacomitinib-d3 (PF-00299804-d3) is the deuterium labeled Dacomitinib. Dacomitinib (PF-00299804) is a specific and irreversible inhibitor of the ERBB family of kinases with IC ₅₀ s of 6 nM, 45.7 nM and 73.7 nM for EGFR, ERBB2, and ERBB4, respectively. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Dacomitinib-d5 (PF-00299804-d5) is the deuterium labeled Dacomitinib. Dacomitinib (PF-00299804) is a specific and irreversible inhibitor of the ERBB family of kinases with IC50 K of 6 nM, 45.7 nM and 73.7 nM for EGFR, ERBB2, and ERBB4, respectively.Purity:>98% Clinical Data:No Development Reported Size:1 mg, 5 mg	

Damnacanthal	Cat. No.: HY-108485	Damnacanthal-d3	Cat. No. : HY-108485S
Damnacanthal is an anthraquinone isolated from the root of Morinda citrifolia. Damnacanthal is a highly potent, selective inhibitor of p56 ^{rck} tyrosine kinase activity.	О О́О́О́Н	Damnacanthal-d3 is the deuterium labeled Damnacanthal. Damnacanthal is an anthraquinone isolated from the root of Morinda citrifolia. Damnacanthal is a highly potent, selective inhibitor of p56 ^{tk} tyrosine kinase activity.	С С С С С С С С С С С С С С С С С С С
Purity: ≥98.0% Clinical Data: No Development Reported Size: 5 mg	o	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	0
Damulin B	Cat. No.: HY-16942	Danshensu (Dan shen suan A; Salvianic acid A)	Cat. No.: HY-N1913
Damulin B is a dammarane-type saponin found in Gynostemma pentaphyllum.Damulin B can induce cell apoptosis and has anti-cancer activities in vitro.	martin ma	Danshensu, an active ingredient of Salvia miltiorrhiza, shows wide cardiovascular benefit by activating Nrf2 signaling pathway.	но с нон
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg	de	Purity: ≥98.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg	
Danivirine		Danivirine-d11	
(TMC120; R147681)	Cat. No.: HY-14266	(TMC120-d11; R147681-d11)	Cat. No.: HY-14266S
Dapivirine (TMC120), the prototype of diarylpyrimidines (DAPY), is an orally active and nonnucleoside reverse transcriptase inhibitor (NRTI). Dapivirine (TMC120) binds directly to HIV-1 reverse transcriptase.		Dapivirine-d11 (TMC120-d11) is the deuterium labeled Dapivirine. Dapivirine (TMC120), the prototype of diarylpyrimidines (DAPY), is an orally active and nonnucleoside reverse transcriptase inhibitor (NRTI) .	
Purity: 99.90% Clinical Data: Phase 3 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	N	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 10 mg	
DAPT		Dasatinib	
(GSI-IX)	Cat. No.: HY-13027	(BMS-354825)	Cat. No.: HY-10181
DAPT (GSI-IX) is a potent and orally active γ -secretase inhibitor with IC ₅₀ s of 115 nM and 200 nM for total amyloid- β (A β) and A β_{42} , respectively. DAPT inhibits the activation of Notch 1 signaling and induces cell differentiation.	F C S H I H S O K	Dasatinib (BMS-354825) is a highly potent, ATP competitive, orally active dual Src/Bcr-Abl inhibitor with potent antitumor activity. The K _i s are 16 pM and 30 pM for Src and Bcr-Abl, respectively.	H010-14-1 H010-14-1 H011-14-1
Purity:99.93%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg		Purity: 99.85% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 200 mg, 500 mg	
Dasatinib hydrochloride (BMS-354825 hydrochloride)	Cat. No. : HY-10181A	Dasatinib monohydrate (BMS-354825 monohydrate)	Cat. No.: HY-10181B
Dasatinib (BMS-354825) hydrochloride is a highly potent, ATP competitive, orally active dual Src/Bcr-Abl inhibitor with potent antitumor activity. The K _s s are 16 pM and 30 pM for Src and Bcr-Abl, respectively.	но-/- ^м О-к <mark>м./</mark> но-/- ^м О-к <mark>м./</mark> н-а	Dasatinib (BMS-354825) monohydrate is a highly potent, ATP competitive, orally active dual Src/Bcr-Abl inhibitor with potent antitumor activity. The K _s are 16 pM and 30 pM for Src and Bcr-Abl, respectively.	Han Han
Purity: 98.86% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 200 mg, 500 mg		Purity:>98%Clinical Data:LaunchedSize:1 mg, 5 mg	

Decetinih-d8		Dauporubicin bydrochlorida (Daupomycin bydroch	Jorido: PD
(BMS-354825-d8)	Cat. No.: HY-10181S	13057 hydrochloride; Rubidomycin hydrochloride)	Cat. No.: HY-13062
Dasatinib D8 is a deuterium labeled Dasatinib. Dasatinib is a dual Bcr-Abl and Src family tyrosine kinase inhibitor. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	HOW NO AND	Daunorubicin (Daunomycin) hydrochloride is a topoisomerase II inhibitor with potent antineoplastic activities. Daunorubicin hydrochloride inhibits DNA and RNA synthesis in sensitive and resistant Ehrlich ascites tumor cells. Purity: 99.23% Clinical Data: Launched Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg	$ \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \end{array}\\ \end{array}\\ \end{array}\\ \end{array} \\ \begin{array}{c} \end{array}\\ \begin{array}{c} \end{array}\\ \end{array} \\ \begin{array}{c} \end{array}\\ \begin{array}{c} \end{array}\\ \end{array} \\ \begin{array}{c} \end{array}\\ \end{array} \\ \begin{array}{c} \end{array}\\ \end{array} \\ \begin{array}{c} \end{array}\\ \begin{array}{c} \end{array}\\ \end{array} \\ \begin{array}{c} \end{array}\\ \begin{array}{c} \end{array}\\ \end{array} \\ \begin{array}{c} \end{array}$ \left(\begin{array}{c} \end{array}\\ \end{array} \\ \left(\begin{array}{c} \end{array}\\ \end{array} \\ \left(\begin{array}{c} \end{array}\\ \end{array}\right) \\ \left(\begin{array}{c} \end{array}\\ \end{array} \\ \left(\begin{array}{c} \end{array}\\ \end{array}\right) \\ \left(\begin{array}{c} \end{array}\\ \end{array} \\ \left(\begin{array}{c} \end{array}\\ \end{array}\right) \\ \left(\begin{array}{c} \end{array}\\ \end{array} \\ \left(\begin{array}{c} \end{array}\\ \end{array}\right) \\ \left(\begin{array}{c} \end{array}\\ \end{array} \\ \left(\begin{array}{c} \end{array}\\ \end{array}\right) \\ \left(\begin{array}{c} \end{array}\\ \left(\begin{array}{c} \end{array}\\ \end{array}\right) \\ \left(\begin{array}{c} \end{array}\\ \left(\begin{array}{c} \end{array}\\ \end{array}) \\ \left(\begin{array}{c} \end{array}\\ \end{array}\right) \\ \left(\begin{array}{c} \end{array}\\ \end{array} \left(\begin{array}{c} \end{array}\\ \left(\begin{array}{c} \end{array}\\ \end{array}) \\ \left(\begin{array}{c} \end{array}\\ \end{array} \left(\begin{array}{c} \end{array} \left(\begin{array}{c} \end{array}\\ \end{array} \left(\begin{array}{c} \end{array}\\ \end{array} \left(\begin{array}{c} \end{array}\\ \end{array} \left(\begin{array}{c} \end{array}\\ \end{array} \left(\begin{array}{c} \end{array} \left(\end{array}) \\ \left(\begin{array}{c} \end{array}\\ \end{array} \left(\end{array}) \\ \left(\end{array} \left) \\ \left(\end{array} \left) \\ \left(\end{array} \left(\end{array} \left) \\ \left) \\ \left(\end{array} \left) \\ \left(\end{array} \left) \\ \left(\end{array} \left) \\ \left) \\
Dauricine	Cat. No.: HY-N0220	DB1976	Cat. No.: HY-135797
Dauricine, a bisbenzylisoquinoline alkaloid in Asiatic Moonseed Rhizome, possesses anti-inflammatory activity. Dauricine inhibits cell proliferation and invasion, and induces apoptosis by suppressing NF-κB activation in a dose- and time-dependent manner in colon cancer. Purity: 99.91%	dente des	DB1976 is a selenophene analog of DB270 and a potent and cell-permeable fully efficacious transcription factor PU.1 inhibitor. Purity: >98%	HAR THE AND
Clinical Data:No Development ReportedSize:5 mg, 10 mg, 20 mg		Clinical Data:No Development ReportedSize:5 mg, 10 mg, 50 mg, 100 mg	
DB1976 dihydrochloride	Cat No . HY-1357974	DB2115 tertahydrochloride	Cat No : HY-124676A
DB1976 dihydrochloride is a selenophene analog of DB270 and a potent and cell-permeable fully efficacious transcription factor PU.1 inhibitor.		DB2115 (tertahydrochloride) is a potent inhibitor of myeloid master regulator PU.1 . DB2115 (tertahydrochloride) has the potential for researching cancers, including hematologic cancers such as leukemia, as well as other conditions	20 ⁴ 0 20
Purity: ≥98.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity: 99.13% Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
DP2212		DPcO	
D5212	Cat. No.: HY-124629	(JRF 12)	Cat. No.: HY-15945
DB2313 is a potent transcription factor PU.1 inhibitor with an apoptosis of 14 nM. DB2313 disrupts the interaction of PU.1 with target gene promoters. DB2313 induces apoptosis of acute myeloid leukemia (AML) cells, and has anticancer effects.	Actor of the	DBeQ is a selective, potent, reversible, and ATP-competitive p97 inhibitor, with an IC ₅₀ value of 1.5 μ M and 1.6 μ M for p97(wt) and p97(C522A), respectively; DBeQ also inhibits Vps4 with an IC ₅₀ of 11.5 μ M.	
Purity:98.13%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg		Purity:99.68%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg	\bigcirc
dRFT6		DRIBR	
	Cat. No.: HY-112588		Cat. No.: HY-117779
dBET6 is a highly potent, selective and cell-permeable PROTAC connected by ligands for Cereblon and BET , with an IC_{50} of 14 nM, and has antitumor activity.	and the second s	DBIBB is a specific nonlipid agonist of the type 2 G protein coupled receptor for lysophosphatidic acid (LPA2). DBIBB mitigates the gastrointestinal radiation syndrome, increases intestinal crypt survival and enterocyte proliferation, and reduces apoptosis .	HO HHY O HY O
Purity: 99.73% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	^{مہ} ی ^ہ ہ ^ہ	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	

DC260126		DC661	
	Cat. No.: HY-101906		Cat. No.: HY-111621
DC260126 is a potent antagonist of GPR40 (FFAR1). DC260126 dose-dependently inhibits GPR40-mediated Ca ²⁺ elevations stimulated by linoleic acid, oleic acid, palmitoleic acid and lauric acid (IC ₅₀ : 6.28, 5.96, 7.07, 4.58 μ M, respectively).	F-C-S-NH O	DC661 is a potent palmitoyl-protein thioesterase 1 (PPT1) inhibitor, inhibits autophagy , and acts as an anti-lysosomal agent. Anti-cancer activity.	Arminet.
Purity: 99.74%		Purity: ≥95.0%	
Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	00 mg	Size: 10 mM \times 1 mL, 2 mg, 5 mg, 10 mg, 25 mg, 50	mg, 100 mg
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		DC70415	
DC1130_00	Cat. No : HV-139108		Cat. No : HV-130603
	Cat. No 111-135100		Cat. No 111-130003
DCH36_06 is a potent and selective p300/CBP inhibitor with IC_{so} of 0.6 μ M and 3.2 μ M for p300 and CBP , respectively. DCH36_06 mediated p300/CBP inhibition leading to hypoacetylation on H3K18 in leukemic cells. Anti-tumor activity.	c The states	DCZ0415, a potent TRIP13 inhibitor, impairs nonhomologous end joining repair and inhibits NF-κB activity. DCZ0415 induces anti-myeloma activity in vitro, in vivo, and in primary cells derived from drug-resistant myeloma patients.	
Purity:99.22%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg		Purity: 99.96% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	00 mg
DC_AC50		Decitabine	
	Cat. No.: HY-107636	(5-Aza-2'-deoxycytidine; 5-AZA-CdR; NSC 127716)	Cat. No.: HY-A0004
DC_AC50 is a dual inhibitor of Atox1 and CCS (copper chaperones). Inhibiting intracellular copper chaperones as a means of reducing/preventing acquired chemotherapy resistance. .		Decitabine (NSC 127716) is an orally active deoxycytidine analogue antimetabolite and a DNA methyltransferase inhibitor.	N N OH
Purity:99.45%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg		Purity: 99.97% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 500 mg, 1 g, 2 g	112N N U
Decursin		Degrasyn	
((+)-Decursin)	Cat. No.: HY-18981	(WP1130)	Cat. No.: HY-13264
Decursin ((+)-Decursin) is a cytotoxic agent and a potent protein kinase C activator from the Root of Angelica gigas. Decursin inhibits tumor growth, migration, and invasion in gastric cancer by down-regulating CXCR7 expression.	of the for	Degrasyn (WP1130) is a cell-permeable deubiquitinase (DUB) inhibitor, directly inhibiting DUB activity of USP9x, USP5, USP14, and UCH37. Degrasyn has been shown to downregulate the antiapoptotic proteins Bcr-Abl and JAK2.	Br N H H
Purity: 99.94% Clinical Data: No Development Reported		Purity: 99.70% Clinical Data: No Development Reported	
Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
Deguelin ((-)-Deguelin; (-)-cis-Deguelin)	Cat. No.: HY-13425	Dehydroaltenusin	Cat. No. : HY-100513A
Deguelin, a naturally occurring rotenoid, acts as a chemopreventive agent by blocking multiple pathways like PI3K-Akt, IKK-NF-ĸB, and MAPK-mTOR-survivin-mediated apoptosis.		Dehydroaltenusin is a small molecule selective inhibitor of eukaryotic DNA polymerase α , a type of antibiotic produced by a fungus with an IC ₅₀ value of 0.68 μ M.	-O C C C C C C C C C C C C C C C C C C C
Purity:99.29%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg	<u>_</u> 0	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	он о



Demethoxycurcumin Demethoxycurcumin-d7 (Curcumin II-d7; Desmethoxycurcumin-d7; (Curcumin II; Desmethoxycurcumin; Monodemethoxycurcumin)Cat. No.: HY-N0006 Cat. No.: HY-N0006S Monodemethoxycurcumin-d7) Demethoxycurcumin(Curcumin II) is a major active Demethoxycurcumin-d7 (Curcumin II-d7) is the curcuminoid: possess anti-inflammatory properties: deuterium labeled Demethoxycurcumin. also exert cytotoxic effects in human cancer cells Demethoxycurcumin(Curcumin II), a major active via induction of apoptosis. IC50 value: Target: in curcuminoid, possess anti-inflammatory properties; also exert cytotoxic effects in human cancer cells vitro: DMC significantly decreased NO secretion by 35-41% in our inflamed cell model. via induction of apoptosis. Purity: >99.0% **Purity:** >98% Clinical Data: No Development Reported Clinical Data: No Development Reported 10 mM × 1 mL, 5 mg, 10 mg, 50 mg Size: Size: 1 mg, 5 mg Demethylzeylasteral Deoxynyboquinone Cat. No.: HY-N0587 Cat. No.: HY-108992 Demethylzeylasteral is a triterpene compound Deoxynyboguinone, an excellent NQO1 substrate, is a potent antineoplastic agent. Deoxynyboquinone isolated from Tripterygium wilfordii Hook F, with anti-inflammatory, immunosuppressive and induces apoptosis in cancer cell lines. anti-tumor activities. Demethylzeylasteral can Deoxynyboquinone kills cancer cells through significantly alleviates atherosclerosis (AS). oxidative stress and reactive oxygen species (ROS) formation. Purity: 99 90% **Purity:** >98% Clinical Data: No Development Reported Clinical Data: No Development Reported 10 mM × 1 mL, 5 mg, 10 mg, 20 mg 1 mg, 5 mg Size: Size: Deracoxib Deoxypodophyllotoxin (SC 046; SC 46; SC 59046) Cat. No.: HY-17509 Cat. No.: HY-N2500 Deoxypodophyllotoxin (DPT), a derivative of Deracoxib, a selective cyclooxygenase-2 inhibitor, podophyllotoxin, is a lignan with potent is a non-narcotic, non-steroidal anti-inflammatory antimitotic, anti-inflammatory and antiviral drug (NSAID). properties isolated from rhizomes of Sinopodophullumhexandrum (Berberidaceae). Purity: 99.86% **Purity:** 99 77% Clinical Data: No Development Reported Clinical Data: Launched Size: 5 mg, 10 mg Size 10 mM × 1 mL, 100 mg, 500 mg Desacetylcinobufotalin Desmethylxanthohumol (Deacetylcinobufotalin) Cat. No.: HY-N0882 Cat. No.: HY-122966

Desacetylcinobufotalin is a natural compound; apoptosis inducer and shows the marked inhibition effect to HepG2 cells and the IC50 value is 0.0279µmol/ml.

99.27% Purity: Clinical Data: No Development Reported Size: 1 mg, 5 mg

Desoxyrhaponticin

Desoxyrhaponticin is a stilbene glycoside from the Tibetan nutritional food Rheum tanguticum Maxim. Desoxyrhaponticin is a Fatty acid synthase (FASN) inhibitor, and has apoptotic effect on human cancer cells.

Purity: 99.80% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg



Cat. No.: HY-N2486

Purity: 99.35% Clinical Data: No Development Reported Size: 1 mg, 5 mg

Metarhizium anisopliae, is one of the

cyclodepsipeptides with insecticidal and

Desmethylxanthohumol is a prenylated

powerful apoptosis inducing agent. Desmethylxanthohumol has antiplasmodial. antiproliferative, and antioxidant bioactivities.

>98%

Purity:

Size

Destruxin B

anticancer activities.

hydroxychalcone isolated from hop cones

Clinical Data: No Development Reported 1 mg, 5 mg

(Humulus lupulus L.). Desmethylxanthohumol is a

Destruxin B, isolated from entomopathogenic fungus



Cat. No.: HY-N6690



Devimistat		Dextran sulfate sodium salt (MW 16000-24000)	
(CPI-613)	Cat. No.: HY-15453		Cat. No.: HY-116282B
Devimistat (CPI-613) is a mitochondrial metabolism inhibitor. Devimistat is a lipoic acid antagonist that abrogates mitochondrial energy metabolism to induce apoptosis in various cancer cells. Purity: 99.59% Clinical Data: Phase 3 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	Green gat	Dextran sulfate sodium salt (MW 16000-24000) is a is a polymer of anhydroglucose with the molecular weight range of 16000-24000. Dextran sulfate sodium salt inhibits the replication of the human immunodeficiency virus by preventing the adsorption of the virus into host cells. Purity: >98% Clinical Data: No Development Reported Size: 100 mg	Dextron sulfate sofure sult (MV 19000-24000)
Dextran sulfate sodium salt (MW 35000-45000)	Cat. No. : HY-116282C	Dextran sulfate sodium salt (MW 4500-5500)	Cat. No.: HY-116282A
Dextran sulfate sodium salt (MW 35000-45000) is a is a polymer of anhydroglucose with the molecular weight range of 35000-45000. Dextran sulfate sodium salt inhibits the replication of the human immunodeficiency virus by preventing the adsorption of the virus into host cells.	Devition Sulfate sodium sati (MM 35000-40900)	Dextran sulfate sodium salt (MW 4500-5500) is a is a polymer of anhydroglucose with the molecular weight range of 4500-5500. Dextran sulfate sodium salt inhibits the replication of the human immunodeficiency virus by preventing the adsorption of the virus into host cells.	Dextran suffate sodium salt (MW 4500-5600)
Clinical Data: No Development Reported Size: 100 mg		Clinical Data: No Development Reported Size: 500 mg	
Dextran sulfate sodium salt (MW 450000-55000	0)	Diallyl Trisulfide	
	Cat. No.: HY-116282D		Cat. No.: HY-117235
Dextran sulfate sodium salt (MW 450000-550000) isa is a polymer of anhydroglucose with themolecular weight range of 450000-550000. Dextransulfate sodium salt inhibits the replication ofthe human immunodeficiency virus by preventing theadsorption of the virus into host cells.Purity:>98%Clinical Data:No Development ReportedSize:100 mg	Denton suffice socium sat (MW 450000-560000)	Diallyl Trisulfide is isolated from Garlic.Diallyl Trisulfide suppresses the growth ofPenicillium expansum (MFC ₉₉ value: ≤ 90 µg/mL) and promotes apoptosis via production ofreactive oxygen species (ROS) and disintegrationof cellular ultrastructure. Anticancer effect.Purity: $\geq 95.0\%$ Clinical Data:No Development ReportedSize:10 mM × 1 mL, 50 mg	s.s.s.
Diatrizoic acid		Diazepinomicin	
(Diatrizoate; Amidotrizoic acid)	Cat. No.: HY-B0926	(ECO-4601; TLN-4601; BU 4664L)	Cat. No.: HY-N6674
Diatrizoic acid (Diatrizoate) is an iodinated radiocontrast agent and has the potential for radiographic imaging of the airways. Diatrizoic acid induces mitochondrial turnover and oxidative stress, and activating apoptosis by dysregulating calcium.Purity:99.86% Clinical Data:No Development Reported Size:10 mM × 1 mL, 500 mg		Diazepinomicin (TLN-4601) is a secondary metabolite produced by Micromonospora sp. Diazepinomicin (TLN-4601) inhibits the EGF-induced Ras-ERK MAPK signaling pathway and induces apoptosis. An anti-tumor agent for K-Ras mutant models. Purity: 98.04% Clinical Data: Phase 2 Size: 1 mg, 5 mg	
Diclofenac	Cat. No.: HY-15036	Diclofenac diethylamine	Cat. No. : HY-15036A
Diclofenac is a potent and nonselective anti-inflammatory agent, acts as a COX inhibitor, with IC ₅₀ s of 4 and 1.3 nM for human COX-1 and COX-2 in CHO cells, and 5.1 and 0.84 μ M for ovine COX-1 and COX-2, respectively.	CI H OH	Diclofenac diethylamine is a potent and nonselective anti-inflammatory agent, acts as a COX inhibitor, with IC ₅₀ s of 4 and 1.3 nM for human COX-1 and COX-2 in CHO cells, and 5.1 and 0.84 μ M for ovine COX-1 and COX-2, respectively.	СІЦОН
Purity: 99.97% Clinical Data: Launched Size: 10 mM × 1 mL, 500 mg, 5 g, 10 g	∽ ci ∽	Purity: 99.93% Clinical Data: Launched Size: 10 mM × 1 mL, 500 mg, 5 g, 10 g	∕ _N ∕

Diclofenac potassium	Cat. No. : HY-15038	Diclofenac Sodium (GP 45840)	Cat. No.: HY-15037
Diclofenac potassium is a potent and nonselective anti-inflammatory agent, acts as a COX inhibitor, with $IC_{so}s$ of 4 and 1.3 nM for human COX-1 and COX-2 in CHO cells, and 5.1 and 0.84 μ M for ovine COX-1 and COX-2, respectively. Purity: \geq 98.0% Clinical Data: Launched Size: 10 mM × 1 mL, 50 mg, 100 mg, 250 mg		Diclofenac Sodium (GP 45840) is a potent and nonselective anti-inflammatory agent, acts as a COX inhibitor, with IC ₅₀ S of 4 and 1.3 nM for human COX-1 and COX-2 in CHO cells, and 5.1 and 0.84 μM for ovine COX-1 and COX-2, respectively. Purity: 99.93% Clinical Data: Launched Size: 10 mM × 1 mL, 500 mg, 5 g	
Diclofenac-13C6 sodium heminonahydrate	Cat. No.: HY-15037S	Diclofenac-d4	Cat. No.: HY-15036S
Diclofenac-13C6 sodium heminonahydrate is the 13C-labeled Diclofenac Sodium. Purity: >98%		Diclofenac-d4 is the deuterium labeled Diclofenac. Diclofenac is a potent and nonselective anti-inflammatory agent, acts as a COX inhibitor, with IC ₅₀ S of 4 and 1.3 nM for human COX-1 and COX-2 in CHO cells, and 5.1 and 0.84 μ M for ovine COX-1 and COX-2, respectively. Purity: >98%	
Clinical Data: No Development Reported Size: 1 mg, 5 mg		Clinical Data: No Development Reported Size: 1 mg, 10 mg	
Diclofenac-d4 sodium	Cat. No.: HY-1503751	Dictamine (Dictamnine; Dectamine)	Cat. No.: HY-N0849
Diclofenac-d4 sodium is the deuterium labeled Diclofenac sodium.		Dictamnine (Dictamine) has the ability to exert cytotoxicity in human cervix, colon, and oral carcinoma cells; A natural plant product has been reported to have antimicrobial activity against bacteria and fungi.	N O
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	D D ONa	Purity:99.10%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg	0
Didesmethylrocaglamide	Cat. No.: HY-19356A	Didymin	Cat. No.: HY-N2068
Didesmethylrocaglamide, a derivative of Rocaglamide, is a potent eukaryotic initiation factor 4A (eIF4A) inhibitor. Didesmethylrocaglamide has potent growth-inhibitory activity with an IC ₅₀ of 5 nM.		Didymin, a dietary flavonoid glycoside from citrus fruits, possesses antioxidant properties. Didymin induces apoptosis by inhibiting N-Myc and upregulating RKIP in neuroblastoma.	
Purity:98.40%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg	O- OH O	Purity:99.90%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 20 mg	
Dienogest (STS 557)	Cat. No.: HY-B0084	Dienogest-d4 (STS 557-d4)	Cat. No.: HY-B0084S
Dienogest(STS-557) is a specific progesterone receptor agonist with potent oral endometrial activity and is used in the treatment of endometriosis. Target: progesterone receptor agonist Dienogest is an orally active synthetic progesterone (or progestin). Purity: 99.83%	O CN O H H H H	Dienogest-d4 is deuterium labeled Dienogest. Purity: >98%	
Clinical Data:LaunchedSize:10 mM × 1 mL, 10 mg, 50 mg, 100 mg		Clinical Data: No Development Reported Size: 1 mg, 5 mg	

Dienogest-d5 (STS 557-d5)	Cat. No. : HY-B0084S1	Dienogest-d6 (STS 557-d6)	Cat. No.: HY-B008452
Dienogest-d5 is deuterium labeled Dienogest.		Dienogest-d6 is deuterium labeled Dienogest.	
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	115) ²
Diffractaic acid	Cat. No.: HY-N2399	Difopein TFA	Cat. No.: HY-P1380A
Diffractaic acid, a major constituent of U. longissimi, acts as an effective proapoptotic agent in various disorders research. Diffractaic acid is the analgesic and antipyretic component of Usnea diffracta.	of of of of	Difopein (TFA), a specific and competitive inhibitor of 14-3-3 protein (a highly conserved eukaryotic regulatory molecule), blocking the ability of 14-3-3 to bind to target proteins and inhibits 14-3-3/Ligand interactions.	
Purity: >98% Clinical Data: No Development Reported Size: 2.5 mg		Purity:94.54%Clinical Data:No Development ReportedSize:1 mg	
Dihydroartemisinin (Dihydroqinghaosu; β-Dihydroartemisinin; Artenimol)	Cat. No.: HY-N0176	Dihydroartemisinin-d3 (Dihydroqinghaosu-d3; β-Dihydroartemisinin-d3; Artenimol-d3)	Cat. No.: HY-N0176S
Dihydroartemisinin is a potent anti-malaria agent.		Dihydroartemisinin-d3 (Dihydroqinghaosu-d3) is the deuterium labeled Dihydroartemisinin. Dihydroartemisinin is a potent anti-malaria agent.	
Clinical Data: Launched Size: 10 mM × 1 mL, 50 mg, 100 mg, 200 mg, 500	mg	Clinical Data: No Development Reported Size: 1 mg, 5 mg	•
Dihydroeponemycin	Cat. No.: HY-108553	Dihydroisotanshinone I	Cat. No.: HY-B1919
Dihydroeponemycin, an analogue of the antitumor and antiangiogenic natural product eponemycin, selectively targets the 20S proteasome .		Dihydroisotanshinone I, a bioactive compound present in danshen, can inhibit the migration of both androgen-dependent and androgen-independent prostate cancer cells. Dihydroisotanshinone I also induces apoptosis and ferroptosis in these lung cancer cells.	
Clinical Data: No Development Reported Size: 1 mg, 5 mg		Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg	
Dihydrokaempferol	Cat. No. : HY-N2897	Dihydrorotenone	Cat. No.: HY-N4202
Dihydrokaempferol is isolated from Bauhinia championii (Benth). Dihydrokaempferol induces apoptosis and inhibits Bcl-2 and Bcl-xL expression. Dihydrokaempferol is a good candidate for new antiarthritic drugs.	HO, C, C, OH OH O	Dihydrorotenone, a natural pesticide, is a potent mitochondrial inhibitor. Dihydrorotenone probably induces Parkinsonian syndrome. Dihydrorotenone induces human plasma cell apoptosis by triggering endoplasmic reticulum stress and activating p38 signaling pathway.	
Purity: 99.88% Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg		Purity: 98.35% Clinical Data: No Development Reported Size: 5 mg, 10 mg	1997-1993

DIM-C-pPhOH		Dinaciclib	
	Cat. No.: HY-112055	(SCH 727965)	Cat. No.: HY-10492
DIM-C-pPhOH is a nuclear receptor 4A1 (NR4A1) antagonist. DIM-C-pPhOH inhibits cancer cell growth and mTOR signaling, induce apoptosis and cellular stress. DIM-C-pPhOH reduces cell proliferation with IC50 values of 13.6 µM and 13.0 µM for ACHN cells and 786-O cells, respectively.	C N OH	Dinaciclib (SCH 727965) is a potent inhibitor of CDK, with IC_{50} s of 1 nM, 1 nM, 3 nM, and 4 nM for CDK2, CDK5, CDK1, and CDK9, respectively.	
Purity: 99.05%		Purity: 99.36%	C
Clinical Data: No Development Reported		Clinical Data: Phase 3	
Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	uu mg	Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
Dinoprost		Dinoprost tromethamine salt (Prostaglandin F2α tr	omethamine
(Prostaglandin F2α; PGF2α)	Cat. No.: HY-12956	salt; PGF2α THAM; Prostaglandin F2α THAM)	Cat. No.: HY-12956A
Dinoprost (Prostaglandin F2α) is an orally active, potent prostaglandin F (PGF) receptor (FP receptor) agonist. Dinoprost is a luteolytic hormone produced locally in the endometrial luminal epithelium and corpus luteum (CL).	На	Dinoprost tromethamine salt (Prostaglandin F2α tromethamine salt) is an orally active, potent prostaglandin F (PGF) receptor (FP receptor) agonist.	но
	(Jan		но он
Purity: 99.06%	но он	Purity: ≥98.0%	HOOH
Clinical Data: Launched		Clinical Data: Launched	00
Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	.00 mg
Dinoprost-d4		Dinoprost-d9	
(Prostaglandin F2a-d4; PGF2α-d4)	Cat. No.: HY-12956S	(Prostaglandin F2a-d9; PGF2α-d9)	Cat. No.: HY-12956S1
Dinoprost-d4 (Prostaglandin F2a-d4) is the deuterium labeled Dinoprost. Dinoprost (Prostaglandin F2 α) is an orally active, potent prostaglandin F (PGF) receptor (FP receptor) agonist.	HQ D D OH	Dinoprost-d9 (Prostaglandin F2a-d9) is the deuterium labeled Dinoprost. Dinoprost (Prostaglandin F2 α) is an orally active, potent prostaglandin F (PGF) receptor (FP receptor) agonist.	но се се
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	но но	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	HO OH D D D D
Disasia		Diagonalia alteracida	
		Diosgenin glucoside	
(Collettiside III; CCRIS 4123)	Cat. No.: HY-N0124		Cat. No.: HY-N0730
Dioscin(CCRIS 4123; Collettiside III) is a natural steroid saponin derived from several plants, showing potent anti-cancer effect against a variety of tumor cell lines.		Diosgenin glucoside, a saponin compound extracted from Tritulus terrestris L., provides neuroprotection by regulating microglial M1 polarization. Diosgenin glucoside protects against spinal cord injury by regulating autophagy and alleviating apoptosis.	
Purity: 99.76%	(2)	Purity: 99.28%	
Clinical Data: Launched		Clinical Data: No Development Reported	
Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg		Size: 5 mg, 10 mg, 20 mg	
Disitertide		Disitertide TFA	
(P144)	Cat. No.: HY-P0118	(P144 TFA)	Cat. No.: HY-P0118A
Disitertide (P144) is a peptidic transforming growth factor-beta 1 (TGF-β1) inhibitor specifically designed to block the interaction with its receptor. Disitertide (P144) is also a PI3K inhibitor and an apoptosis inducer. br/>.	TSLDASIIWAMMQN	Disitertide (P144) TFA is a peptidic transforming growth factor-beta 1 (TGF- β 1) inhibitor specifically designed to block the interaction with its receptor. Disitertide (P144) TFA is also a P13K inhibitor and an apoptosis inducer. br/>.	TSLDASIIWAMMON (TFA sait)
Purity: >98%		Purity: 95.87%	
Clinical Data: Phase 2		Clinical Data: Phase 2	
Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg		Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg	

DL-Cystathionine dihydrochloride DJ001 Cat. No.: HY-133146 Cat. No.: HY-W009749B DJ001 is a highly specific, selective and DL-Cystathionine dihydrochloride is a racemic non-competitive protein tyrosine phosphatase- σ melange of the L-Cystathionine dihydrochloride and D-Cystathionine dihydrochloride. L-Cystathionine (PTP\sigma) inhibitor with an IC $_{\rm 50}$ of 1.43 $\mu M.$ DJ001 dihydrochloride is a nonprotein thioether and is a displays no inhibitory activity against other key amino acid associated with the metabolic state phosphatases, with only modest inhibitory activity against Protein Phosphatase 5. of sulfur-containing amino acids. Purity: 99.59% Purity: >98% Clinical Data: No Development Reported Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg Size: 1 mg, 5 mg dMCL1-2 DMH2 Cat. No.: HY-128360 (VU364849) Cat. No.: HY-110245 dMCL1-2 is a potent and selective PROTAC of DMH2 is a potent BMP receptor antagonist. DMH2 myeloid cell leukemia 1 (MCL1) (Bcl-2 family downregulates the expression of Id1 and Id3 member) based on Cereblon, which binds to MCL1 with proteins, and inhibits the proliferation and a K_p of 30 nM. dMCL1-2 activats the cellular induces cell death of lung cancer cell lines. apoptosis machinery by degradation of MCL1. Purity: > 98% **Purity:** >98% Clinical Data: No Development Reported Clinical Data: No Development Reported Size: 1 mg, 5 mg Size: 1 mg, 5 mg DMU-212 DMUP Cat. No.: HY-137977 Cat. No.: HY-115983 DMUP is a potent CD47-SIRPa axis inhibitor. DMUP DMU-212 is a methylated derivative of Resveratrol (HY-16561), with antimitotic, anti-proliferative, induces apoptosis and increases the macrophage antioxidant and apoptosis promoting activities. phagocytosis in A549 cells. DMUP decreases the DMU-212 induces mitotic arrest via induction of expression of CD47 and SIRPα protein. DMUP shows apoptosis and activation of ERK1/2 protein. antitumor activity. DMU-212 has orally active. 99.91% Purity: >98% Purity: Clinical Data: No Development Reported Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 25 mg, 50 mg Size 1 mg, 5 mg Dobesilate-d6 calcium Docetaxel (RP-56976) Cat. No.: HY-111603S Cat. No.: HY-B0011 Docetaxel (RP-56976) is a Dobesilate-d6 (calcium) is deuterium labeled Calcium dobesilate. Calcium dobesilate, a microtubule depolymerization inhibitor, vasoprotective, is widely used in chronic venous with an IC_{so} of 0.2 μ M. Docetaxel attenuates the disease, diabetic retinopathy and the symptoms of effects of bcl-2 and bcl-xL gene expression. haemorrhoidal attack in many countries. Docetaxel arrests the cell cycle at G2/M and leads to cell apoptosis. >98% Purity: 99.96% Purity: Clinical Data: No Development Reported Clinical Data: Launched Size: 1 mg, 5 mg Size 10 mM × 1 mL, 100 mg, 200 mg **Docetaxel Trihydrate** Docetaxel-d5 trihydrate (RP-56976 Trihydrate) Cat. No.: HY-B0011A (RP-56976-d5 trihydrate) Cat. No.: HY-B0011AS Docetaxel Trihydrate (RP-56976 Trihydrate) is an Docetaxel-d5 (RP-56976-d5) trihydrate is the antineoplastic agent and inhibits deuterium labeled Docetaxel (Trihydrate). microtubule depolymerization with an IC₅₀ Docetaxel Trihydrate (RP-56976 Trihydrate) is an value of 0.2 μ M. Docetaxel Trihydrate is a antineoplastic agent and inhibits semisynthetic analog of taxol and attenuates the. microtubule depolymerization with an IC₅₀ value of 0.2 µM. 99.92% **Purity:** >98% Purity: Clinical Data: Launched Clinical Data: No Development Reported Size: 10 mM × 1 mL, 100 mg Size: 1 mg, 5 mg

Docetaxel-d9 (RP-56976-d9)	Cat. No. : HY-B0011S	Dolastatin 15 (DLS 15)	Cat. No.: HY-P1126
$\begin{array}{llllllllllllllllllllllllllllllllllll$		Dolastatin 15 (DLS 15), a depsipeptide derived from Dolabella auricularia, is a potent antimitotic agent structurally related to the antitubulin agent Dolastatin 10. Dolastatin 15 induces cell cycle arrest and apoptosis in multiple myeloma cells.Purity:>98% Clinical Data:Clinical Data:No Development Reported 	
Domatinostat (4SC-202 free base)	Cat. No.: HY-16012A	Domatinostat tosylate (4SC-202)	Cat. No.: HY-16012
Domatinostat (4SC-202 free base) is a selective class I HDAC inhibitor with IC ₅₀ of 1.20 μ M, 1.12 μ M, and 0.57 μ M for HDAC1, HDAC2, and HDAC3, respectively. It also displays inhibitory activity against Lysine specific demethylase 1 (LSD1).	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Domatinostat tosylate (4SC-202) is a selective class I HDAC inhibitor with IC ₅₀ of 1.20 μ M, 1.12 μ M, and 0.57 μ M for HDAC1, HDAC2, and HDAC3, respectively. It also displays inhibitory activity against Lysine specific demethylase 1 (LSD1).	affrantio a
Purity: 99.08% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity: 99.66% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
Doxorubicin (Hydroxydaunorubicin)	Cat. No.: HY-15142A	Doxorubicin hydrochloride (Hydroxydaunorubicin hydrochloride)	Cat. No. : HY-15142
Doxorubicin (Hydroxydaunorubicin), a cytotoxic anthracycline antibiotic, is an anti-cancer chemotherapy agent. Doxorubicin inhibits topoisomerase II with an IC_{50} of 2.67 μ M, thus stopping DNA replication.		Doxorubicin (Hydroxydaunorubicin) hydrochloride, a cytotoxic anthracycline antibiotic, is an anti-cancer chemotherapy agent. Doxorubicin hydrochloride is a potent human DNA topoisomerase I and topoisomerase II inhibitor with IC _{so} s of 0.8 µM and 2.67 µM, respectively.	
Purity: >98% Clinical Data: Launched Size: 5 mg, 10 mg, 25 mg		Purity: 99.47% Clinical Data: Launched Size: 10 mM × 1 mL, 50 mg, 100 mg, 200 mg, 500 mg/state	ng, 1 g
Dp44mT		ррво	
	Cat. No.: HY-18973		Cat. No.: HY-U00441
Dp44mT is an iron chelator with selective anticancer activity.		DPBQ activates p53 and triggers apoptosis in a polyploid-specific manner, but does not inhibit topoisomerase or bind DNA. DPBQ elicits expression and phosphorylation of p53 and this effect is specific to tetraploid cells.	
Purity: ≥98.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 25 mg, 50 mg, 100 mg	N S	Purity: ≥98.0% Clinical Data: No Development Reported Size: 5 mg	· · ·
DPN (Diarylpropionitrile)	Cat. No.: HY-12452	Dracorhodin perchlorate	Cat. No.: HY-N0726
DPN (Diarylpropionitrile) is a non-steroidal estrogen receptor β (ER β) selective ligand, with an EC ₅₀ of 0.85 nM. DPN has neuroprotective effects in a number of neurological diseases.	но	Dracorhodin perchlorate (Dracohodin perochlorate) is a natural product extracted from a natural medicine Dragon's blood. Dracorhodin perchlorate inhibits cell proliferation and induces cell cycle arrest and apoptosis.	
Purity:99.66%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 500 mg		Purity:98.45%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 20 mg	

Drolovifene		Drovinostat	
(3-Hydroxytamoxifen)	Cat. No.: HY-121149	(NS 41080)	Cat. No.: HY-13267
Droloxifene, a Tamoxifen derivative, is an orally active and selective estrogen receptor modulator. Droloxifene shows antiestrogenic and anti-implantation effects. Droloxifene induces p53 expression and apoptosis in MCF-7 cells. Purity: 99.68% Clinical Data: No Development Reported Size: 5 mg	N-0 CT	Droxinostat(NS41080) is a selective inhibitor of HDAC3, HDAC6, and HDAC8 with IC50 of 16.9, 2.47 and 1.46 µM, respectively; > 8-fold selective against HDAC3 and no inhibition to HDAC1, 2, 4, 5, 7, 9, and 10. Purity: 99.60% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	a for the second
Dubermatinib (TP-0903)	Cat. No.: HY-12963	Duocarmycin A	Cat. No.: HY-12455
Dubermatinib (TP-0903) is a potent and selective Axl receptor tyrosine kinase inhibitor with an IC_{s0} value of 27 nM.		Duocarmycin A, which is one of well-known antitumor antibiotics, is a DNA alkylator and efficiently alkylates adenine N3 at the 3' end of AT-rich sequences in the DNA.	H H H NH NH OF NH O
Purity: 99.82% Clinical Data: Phase 1 Size: 5 mg, 10 mg, 50 mg, 100 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	4°
DuP-697	Cat. No.: HY-103387	Dutasteride (GG 745; GI 198745)	Cat. No.: HY-13613
DuP-697 is a member of the vicinal diaryl heterocycles and a potent, irreversible, selective and orally active COX-2 inhibitor (IC_{s0} of 10 nM and 800 nM for human COX-2 and COX-1, respectively).	Br S	Dutasteride (GG745) is a potent inhibitor of both 5α -reductase isozymes. Dutasteride may possess off-target effects on the androgen receptor (AR) due to its structural similarity to DHT.	
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	F	Purity: 99.75% Clinical Data: Launched Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg	o≪ N H
F64FC26		FAD1	
	Cat. No.: HY-122895		Cat. No.: HY-123056
E64FC26 is a potent pan-inhibitor of the protein disulfide isomerase (PDI) family, with IC_{50} s of 1.9, 20.9, 25.9, 16.3, and 25.4 µM against PDIA1, PDIA3, PDIA4, TXNDC5, and PDIA6, respectively. E64FC26 shows anti-myeloma activity.	б сон	EAD1 is a potent autophagy inhibitor with antiproliferative activity in lung and pancreatic cancer cells. EAD1 also induces apoptosis .	janne.
Purity: 99.37% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	0 mg	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
EB-3D	Cat. No.: HY-115463	EC359	Cat. No.: HY-120142
EB-3D is a potent and selective choline kinase α (ChoK α) inhibitor, with an IC ₅₀ of 1 μ M for ChoK α 1. EB-3D exerts effects on ChoK α expression, AMPK activation, apoptosis , endoplasmic reticulum stress and lipid metabolism.	~q.0~~0~Q.	EC359 is a potent, selective, high affinity and orally active leukemia inhibitory factor receptor (LIFR) inhibitor with a K _a of 10.2 nM, which directly interacts with LIFR to effectively block LIF/LIFR interactions.	
Purity:98.78%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg		Purity:98.11%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	



EGFR-IN-45	Cat. No : HV-145867	EGFR-IN-46	Cat No . HV-144794
$\begin{array}{llllllllllllllllllllllllllllllllllll$		EGFR-IN-46 is a potent EGFR and FAK dual inhibitor with IC ₅₀ s of 20.17 nM, 14.25 nM, respectively. EGFR-IN-46 significantly inhibits the growth of cancer cells. EGFR-IN-46 induces cell apoptosis. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	FO ^L Contraction
EGFR-IN-47	Cat. No.: HY-143337	ЕІ1 (КВ-145943)	Cat. No.: HY-15573
EGFR-IN-47 is a potent and orally active EGFR ^{LSSB/T790M/CP975} inhibitor with an IC_{s0} of 0.01 μ M. EGFR-IN-47 induces cell cycle attest and cell apoptosis . EGFR-IN-47 has the potential for the research of NSCLC.		EI1 (KB-145943) is a potent and selective EZH2 inhibitor with IC_{so} of 15 nM and 13 nM for EZH2 (WT) and EZH2 (Y641F), respectively.	
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	ν, γ	Purity:99.18%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	2
EL-102	Cat. No.: HY-16187	Elesclomol (STA-4783)	Cat. No.: HY-12040
EL102 is a inhibitor of HIF1 α , Which can inhibit tubulin polymerisation and decreased microtubule stability. target: HIF1 α IC 5020-40 nM.	N=-() () () () () () () () () () () () () (Elesclomol (STA-4783) is an oxidative stress inducer that induces cancer cell apoptosis . Elesclomol is a reactive oxygen species (ROS) inducer. Elesclomol shows antitumor activity against a broad range of cancer cell types.	
Purity: 99.76% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 50 mg, 100	mg	Purity: 99.80% Clinical Data: Phase 3 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
ELR510444	Cat. No. : HY-16191	Emamectin Benzoate (MK-244)	Cat. No. : HY-B0837
ELR510444 is a novel microtubule disruptor; inhibits MDA-MB-231 cell proliferation with IC50 of 30.9 nM; not a substrate for the P-glycoprotein drug transporter and retains activity in β III-tubulin-overexpressing cell lines.Purity: \geq 98.0% Clinical Data: No Development Reported Size:10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	NB-S-CT95CT	Emamectin Benzoate (MK-244) is an orally active nervoussystem toxicant by binding g-aminobutyric (GABA) receptor in insects. Emamectin Benzoate is one of semi-synthetic derivative of Avermectin (HY-15311) with a broadspectrum of insecticidal and acaricidal activity. Purity: 99.40% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 100 mg, 500 mg	or the statest of
Embelin (Embelic acid; Emberine; NSC 91874)	Cat. No.: HY-17473	ENMD-2076	Cat. No.: HY-10987A
Embelin (Embelic acid), a potent, nonpeptidic XIAP inhibitor (IC_{s0} =4.1 µM), inhibits cell growth, induces apoptosis , and activates caspase-9 in prostate cancer cells with high levels of XIAP.	но в	ENMD-2076 is a multi-targeted kinase inhibitor with IC _{so} s of 1.86, 14, 58.2, 15.9, 92.7, 70.8, 56.4 nM for Aurora A, Flt3, KDR/VEGFR2, Flt4/VEGFR3, FGFR1, FGFR2, Src, PDGFRα, respectively.	
Purity:98.75%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg, 50 mg		Purity: 99.12% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	Ų
ENMD-2076 Tartrate		Enniatin A1	
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	Cat. No.: HY-10987		Cat. No.: HY-N6704
ENMD-2076 Tartrate is a multi-targeted kinase inhibitor with IC _{so} s of 1.86, 14, 58.2, 15.9, 92.7, 70.8, 56.4 nM for Aurora A , Flt3 , KDR/VEGFR2, Flt4/VEGFR3, FGFR1, FGFR2, Src, PDGFRα, respectively.		Enniatin A1 isolated from Fusarium mycotoxins is a cyclic hexadepsipeptide consisting of alternating D- α -hydroxyisovaleric acids and N-methyl-L-amino acids.	
Purity: 98.87% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 5 mg, 10 mg		Purity:>98%Clinical Data:No Development ReportedSize:5 mg	L º Y
Enniatin complex		Enterodiol	
	Cat. No.: HY-N6706		Cat. No.: HY-108695
Enniatin complex is a mixture of cyclohexadepsipeptides isolated largely from Fusarium species of fungi, and has ionophoric, antibiotic, and in vitro hypolipidaemic properties.	Enniatin complex	Enterodiol is transformed by human intestinal bacteria from lignans contained in various whole-grain cereals, nuts, legumes, flaxseed, and vegetables. Enterodiol has an apoptotic effect in colorectal cancer (CRC) cells. Anti-cancer activities.	HO. HO.
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
Enterolactone		Enterolactone-d6	
	Cat. No.: HY-108692		Cat. No.: HY-108692S
Enterolactone is a bioactive phenolic metabolite known as a mammalian lignan derived from dietary lignans. Enterolactone has estrogenic properties and anti-breast cancer activity.	ОН	Enterolactone-d6 is the deuterium labeled Enterolactone. Enterolactone is a bioactive phenolic metabolite known as a mammalian lignan derived from dietary lignans. Enterolactone has estrogenic properties and anti-breast cancer activity.	HO D OF CH
Purity: >98% Clinical Data: No Development Reported Size: 500 μg	ОН	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
Entinostat		Enzastaurin	
(MS-275; SNDX-275)	Cat. No.: HY-12163	(LY317615)	Cat. No.: HY-10342
Entinostat is an oral and selective class I HDAC inhibitor, with IC_{so} s of 243 nM, 453 nM, and 248 nM for HDAC1, HDAC2, and HDAC3, respectively.	Collectore Hand	Enzastaurin (LY317615) is a potent and selective $PKC\beta$ inhibitor with an IC_{s_0} of 6 nM, showing 6-to 20-fold selectivity over PKC α , PKC γ and PKC ϵ .	
Purity: 99.65% Clinical Data: Phase 3 Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg]	Purity: 99.92% Clinical Data: Phase 3 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg,	200 mg
EPI-001	Cat. No. : HY-100348	Epibrassinolide (24-Epibrassinolide; B1105; BP55)	Cat. No.: HY-N0848
EPI-001, a selective inhibitor of Androgen Receptor (AR) , targets transactivation unit 5 (Tau-5) of the AR. EPI-001 can inhibit transactivation of the AR amino-terminal domain (NTD), with an IC ₅₀ of ~6 μ M. EPI-001 is also a selective modulator of PPARy .	но то сто сто сто	Epibrassinolide (24-Epibrassinolide) is a ubiquitously occurring plant growth hormone which shows great potential to alleviate heavy metals and pesticide stress in plants. Epibrassinolide is a potential apoptotic inducer in various cancer cells without affecting the non-tumor cell growth.	
Purity:98.52%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg, 25 mg, 50 mg		Purity: ≥98.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg,	500 mg





Estramustine phosphate sodium	Cat. No.: HY-13627	Etalocib (LY293111; VML 295)	Cat. No.: HY-13628
Estramustine phosphate sodium, an estradiol analog, is an orally active antimicrotubule chemotherapy agent. Estramustine phosphate sodium depolymerises microtubules by binding to microtubule associated proteins (MAPs) and/or to tubulin. Purity: 99.42% Clinical Data: Launched Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg		Etalocib (LY293111), an orally active leukotriene B_4 receptor antagonist, inhibits the binding of $[^{1}H]LTB_4$, with a K ₁ of 25 nM. Etalocib(LY293111) prevents LTB ₄ -induced calciummobilization with an IC ₅₀ of 20 nM. Etalocib(LY293111) induces apoptosis.Purity:98.27%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	2°°, 2°°, 2°°, 2°°,
Ethoxysanguinarine	Cat. No.: HY-N4317	Ethyl 3,4-dihydroxybenzoate (Ethyl protocatechuate)	Cat. No.: HY-W016409
Ethoxysanguinarine is a benzophenanthridine alkaloid natural product that is mainly found in Macleaya cordata. Ethoxysanguinarine inhibits viability and induces apoptosis of colorectal cancer cells by inhibiting protein phosphatase 2A (CIP2A). Purity: 99.73% Clinical Data: No Development Reported Size: 5 mg, 10 mg		Ethyl 3,4-dihydroxybenzoate (Ethyl protocatechuate), an antioxidant, is a prolyl-hydroxylase inhibitor found in the testa of peanut seeds. Ethyl 3,4-dihydroxybenzoate protects myocardium by activating NO synthase and generating mitochondrial ROS.Purity:99.85%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 500 mg	HOHO
Ethylene dimethanesulfonate	Cat. No.: HY-129524	Etidronic acid (Etidronate; HEDPA; HEDP)	Cat. No.: HY-B0302
Ethylene dimethane sulfonate is a mild alkylating, non-volatile methanesulfonic diester of ethylene glycol. Ethylene dimethanesulfonate has selective pro-apoptotic effects on LCs.		Etidronic acid (Etidronate) is a bisphosphonate used in detergents, water treatment, cosmetics and pharmaceutical treatment.	НО. Р≠О 0 Р≠О НО-Р
Purity: ≥98.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 25 mg, 50 mg, 100 mg		Purity: ≥98.0% Clinical Data: Launched Size: 10 mM × 1 mL, 500 mg, 1 g	НО ОН
Etomovir		Etomovir codium colt	
((R)-(+)-Etomoxir)	Cat. No.: HY-50202	((R)-(+)-Etomoxir sodium salt)	Cat. No.: HY-50202A
Etomoxir ((R)-(+)-Etomoxir) is an irreversible inhibitor of carnitine palmitoyltransferase 1a (CPT-1a), inhibits fatty acid oxidation (FAO) through CPT-1a and inhibits palmitate β -oxidation in human, rat and guinea pig.	°O.~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Etomoxir((R)-(+)-Etomoxir) sodium salt is an irreversible inhibitor of carnitine palmitoyltransferase 1a (CPT-1a), inhibits fatty acid oxidation (FAO) through CPT-1a and inhibits palmitate β -oxidation in human, rat and guinea pig.	al Constant of the second seco
Purity:99.92%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg		Purity:99.46%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg	
Etoposide (VP-16; VP-16-213)	Cat. No.: HY-13629	Etoposide phosphate (BMY-40481)	Cat. No. : HY-13630
Etoposide (VP-16; VP-16-213) is an anti-cancer chemotherapy agent. Etoposide inhibits topoisomerase II , thus stopping DNA replication. Etoposide induces cell cycle arrest, apoptosis and autophagy .		Etoposide phosphate (BMY-40481) is a potent anti-cancer chemotherapy agent and a selective topoisomerase II inhibitor to prevent re-ligation of DNA strands.	
Purity: 99.94% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 200 mg, 500 mg	ò-(° () он	Purity: 98.40% Clinical Data: Launched Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg,	

Etoposide phosphate disodium		Etoposide-13C,d3	
(BMY-40481 disodium)	Cat. No.: HY-13630A	(VP-16-13C,d3; VP-16-213-13C,d3)	Cat. No.: HY-13629S1
Etoposide phosphate disodium (BMY-40481 disodium) is a potent anti-cancer chemotherapy agent and a selective topoisomerase II inhibitor to prevent re-ligation of DNA strands. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Etoposide-13C,d3 is the 13C- and deuterium labeled. Etoposide (VP-16; VP-16-213) is an anti-cancer chemotherapy agent. Etoposide inhibits topoisomerase II, thus stopping DNA replication. Etoposide induces cell cycle arrest, apoptosis and autophagy. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	$H_{0} \xrightarrow{I_{0}}_{I_{0}} \xrightarrow{I_{0}}_{I_{1}} H_{0} \xrightarrow{I_{0}}_{I_{0}} \xrightarrow{I_{0}}_$
Ftretinate		Etretinate-d3	
(Ro 10-9359)	Cat. No.: HY-B0797		Cat. No.: HY-B0797S
Etretinate(Ro 10-9359) is a second-generation retinoid that has the potential for severe psoriasis treatment.	Julia	Etretinate-d3 is the deuterium labeled Etretinate. Etretinate (Ro 10-9359) is a second-generation retinoid that has the potential for severe psoriasis research.	statestin
Purity: 98.04% Clinical Data: Launched Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg		Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 10 mg	
Eugenol		Eugenol-d3	
Euganal is an assential ail found in claues with	Cat. No.: HY-N0337	Euganol d2 is the douterium labeled Euganol	Cat. No.: HY-N0337S
antibacterial, anthelmintic and antioxidant activity. Eugenol is shown to inhibit lipid peroxidation.	HO	Eugenol-as is the deuterium labeled Eugenol. Eugenol is an essential oil found in cloves with antibacterial, anthelmintic and antioxidant activity. Eugenol is shown to inhibit lipid peroxidation.	OHD D
Purity: 98.45% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 500 mg		Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 50 mg	
Eupalinolide O	Cat. No.: HY-N8187	Euphorbia Factor L1	Cat. No.: HY-N2557
Eupalinolide O is a sesquiterpene lactone with anticancer activities. Eupalinolide O induces cell apoptosis in human MDA-MB-468 breast cancer cells.		Euphorbia Factor L1 is a diterpenoid from Euphorbia lathyris L., reduces the expression of Bcl-2, PI3K, AKT and mTOR protein and mRNA, upregulates cleaved caspase-9 and caspase-3 levels, buts shows no effect on pro-caspase-9 and pro-caspase-3.	
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg	Ö	Purity:99.69%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 20 mg	
Euphorbia Factor L2	Cat. No.: HY-N5001	Euscaphic acid	Cat. No.: HY-N2566
Euphorbia factor L2, a lathyrane diterpenoid isolated from caper euphorbia seed (the seeds of Euphorbia lathyris L.), has been traditionally applied to treat cancer. Euphorbia factor L2 shows potent cytotoxicity and induces apoptosis via a mitochondrial pathway. Purity: 99.65% Clinical Data: No Development Reported		Euscaphic acid, a DNA polymerase inhibitor, is a triterpene from the root of the R. alceaefolius Poir. Euscaphic inhibits calf DNA polymerase α (pol α) and rat DNA polymerase β (pol β) with IC ₅₀ values of 61 and 108 μ M. Euscaphic acid induces apoptosis . Purity: 98.34% Clinical Data: No Development Reported	
Size: 10 mM × 1 mL, 5 mg, 10 mg		Size: 1 mg, 5 mg, 10 mg	



Falcarindiol	Cat. No. : HY-N0364	Famitinib (SHR1020)	Cat. No.: HY-108713
Falcarindiol, an orally active polyacetylenic oxylipin, activates PPARy and increases the expression of the cholesterol transporter ABCA1 in cells. Falcarindiol induces apoptosis and autophagy .	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Famitinib (SHR1020), an orally active multi-targeted kinase inhibitor, inhibits the activity of c-kit, VEGFR-2 and PDGFR β with IC _{s0} values of 2.3 nM, 4.7 nM and 6.6 nM, respectively.	
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg		Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
Fangchinoline	Cat. No.: HY-N1372A	Farudodstat (ASLAN003)	Cat. No.: HY-129239
Fangchinoline is isolated from Stephania tetrandra with extensive biological activities, such as enhancing immunity, anti-inflammatory sterilization and anti-atherosclerosis.		Farudodstat (ASLAN003) is an orally active and potent Dihydroorotate Dehydrogenase (DHODH) inhibitor with an IC_{s0} of 35 nM for human DHODH enzyme. Farudodstat inhibits protein synthesis via activation of AP-1 transcription factors.	
Purity:99.92%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg	L _N H	Purity: 99.95% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	00 mg
Fascaplysin	Cat. No.: HY-112328	FB23-2	Cat. No.: HY-127103
Fascaplysin is an antimicrobial and cytotoxic red pigment, that can come from the marine sponge (Fascaplysinopsis sp.). Fascaplysin has been synthesized in seven steps from indole (65% yield).		FB23-2 is a potent and selective inhibitor of mRNA N ⁶ -methyladenosine (m ⁶ A) demethylase FTO , with an IC ₅₀ of 2.6 μ M. FB23-2 has anti-proliferation activity. FB23-2 can be used for the research of acute myeloid leukemia (AML).	
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Purity:99.53%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg	0~
FD223	Cat. No.: HY-132231	Fedratinib (TG-101348; SAR 302503)	Cat. No.: HY-10409
FD223 is a potent and selective phosphoinositide 3-kinase delta (PI3Kδ) inhibitor. FD223 displays high potency (IC _{s0} =1 nM) and good selectivity over other isoforms (IC _{s0} s of 51 nM, 29 nM and 37 nM, respectively for α, β and γ).	C C C C C C C C C C C C C C C C C C C	Fedratinib (TG-101348) is a potent, selective, ATP-competitive and orally active JAK2 inhibitor with IC_{so}^{5} of 3 nM for both JAK2 and JAK2V617F kinase. Fedratinib shows 35- and 334-fold selectivity over JAK1 and JAK3, respectively.	
Purity:98.68%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg		Purity: 99.87% Clinical Data: Launched Size: 10 mM × 1 mL, 5 mg, 10 mg, 100 mg, 200 mg,	500 mg, 1 g
Fedratinib hydrochloride hydrate (TG-101348 hydrod hydrate; SAR 302503 hydrochloride hydrate)	chloride Cat. No.: HY-10409A	Fenobucarb	Cat. No.: HY-B0835
Fedratinib hydrochloride hydrate (TG-101348 hydrochloride hydrate) is a potent, selective, ATP-competitive and orally active JAK2 inhibitor with IC ₅₀ s of 3 nM for both JAK2 and JAK2V617F kinase.		Fenobucarb is a carbamate insecticide. Fenobucarb induces zebrafish developmental neurotoxicity through pathways involved in inflammation, oxidative stress, degeneration and apoptosis.	H o t
Purity: 99.86% Clinical Data: Launched Size: 10 mM × 1 mL, 5 mg, 10 mg, 100 mg, 200 mg, 5	н-а 500 mg, 1 g	Purity:99.60%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 250 mg	0 📡

Fenobucarb-d3	Cat No · HY-B0835S	Fenoprofen Calcium hydrate	Cat No : HY-B0288B
Fenobucarb-d3 is the deuterium labeled Fenobucarb.Fenobucarb is a carbamate insecticide. Fenobucarbinduces zebrafish developmental neurotoxicitythrough pathways involved in inflammation,oxidative stress, degeneration and apoptosis.Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Fenoprofen Calcium hydrate is a nonsteroidal, anti-inflammatory antiarthritic agent. Purity: 99.93% Clinical Data: Launched Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g	0.5Ca ²⁺
FeTPPS	Cat. No.: HY-131697	FGFR4-IN-7	Cat. No.: HY-115902
FeTPPS, a 5,10,15,20-tetrakis (4-sulfonatophenyl) porphyrin iron III chloride peroxynitrite decomposition catalyst, possesses evident neuroprotective effects in a experimental model of spinal cord damage. FeTPPS acts as a. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.0	FGFR4-IN-7 (Compound C3) is a covalent reversible FGFR4 inhibitor with an IC ₅₀ value of 0.42 µM. FGFR4-IN-7 induces apoptosis via the FGFR4 signaling pathway blockage. FGFR4-IN-7 can be used for the research of hepatocellular carcinoma (HCC). Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	∼ ¹ ₆ k∽k ^t tr _o ço.
Fidaxomicin		Fidaxomicin-d7	
(OPT-80; PAR-101) Fidaxomicin (OPT-80), a macrocyclic RNA polymerase inhibitor, has a narrow spectrum of activity. Fidaxomicin selectively eradicates pathogenic Clostridium difficile with minimal disruption to the multiple species of bacteria that make up the normal, healthy intestinal flora. Purity: 99.85% Clinical Data: Launched Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	Cat. No.: HY-17580	Fidaxomicin-D7 (OPT-80-D7) is the deuterium labeled Fidaxomicin. Fidaxomicin (OPT-80), a macrocyclic RNA polymerase inhibitor, has a narrow spectrum of activity. Purity: >98% Clinical Data: No Development Reported Size: S00 µg, 5 mg, 25 mg	Cat. No.: HY-17580S
Filanesib (ARRY-520)	Cat. No.: HY-15187	Fimasartan (BR-A-657)	Cat. No.: HY-B0780
Filanesib (ARRY-520) is a selective and noncompetitive kinesin spindle protein (KSP) inhibitor, with an IC ₅₀ of 6 nM for human KSP. Filanesib induces cell death by apoptosis in vitro. Filanesib has potent anti-proliferative activity. Purity: 99.59% Clinical Data: Phase 1 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Fimasartan(BR-A-657) is a non-peptide angiotensin II receptor antagonist used for the treatment of hypertension and heart failure. Purity: 98.04% Clinical Data: Launched Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
Fimasartan-d6 (BR-A-657-d6)	Cat. No.: HY-B0780S	Fimepinostat (CUDC-907)	Cat. No.: HY-13522
Fimasartan-d6 is deuterium labeled Fimasartan.		Fimepinostat (CUDC-907) potently inhibits class I PI3Ks as well as classes I and II HDAC enzymes with an IC ₅₀ of 19/54/39 nM and 1.7/5.0/1.8/2.8 nM for PI3K α /PI3K β /PI3K δ and HDAC1/HDAC2/HDAC3/HDAC10, respectively.	
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	a☆a	Purity: 99.95% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	

Flavokawain A		Flavokawain B	
	Cat. No.: HY-N2420	(Flavokavain B)	Cat. No.: HY-N2132
Flavokawain A, a proming anticarcinogenic agent , is a chalcone from kava extract with anti-tumor activity. Flavokawain A induces cell apoptosis by involvement of Bax protein-dependent and mitochondria-dependent apoptotic pathway.	of the of	Flavokawain B (Flavokavain B) is a chalcone isolated from the root extracts of kava-kava plant and a potent apoptosis inducer for inhibiting the growth of various cancer cell lines. Flavokawain B (Flavokavain B) shows strong antiangiogenic activity.	
Purity:99.93%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 50 mg		Purity:99.90%Clinical Data:No Development ReportedSize:10 mg	
Flavokawain C	Cat. No.: HY-N2445	Flavopiridol (HMR-1275; Alvocidib; L86-8275)	Cat. No.: HY-10005
Flavokawain C is a natural chalcone found in Kava root. Flavokawain C exerts cytotoxicity against human cancer cell lines, with an IC_{s0} of 12.75 μ M for HCT 116 cells.	OH O E	Flavopiridol (Alvocidib) is a broad spectrum and competitive inhibitor of CDKs , inhibiting CDK1, CDK2, CDK4 with $IC_{so}s$ of 30, 170, 100 nM, respectively.	HO HO CI
Purity:≥98.0%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 20 mg		Purity: 99.72% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	Ņ
FLLL32		Floxuridine	
	Cat. No.: HY-100544	(5-Fluorouracil 2'-deoxyriboside)	Cat. No.: HY-B0097
FLLL32, a synthetic analog of curcumina, is a JAK2/STAT3 dual inhibitor with anti-tumor activity. FLLL32 can inhibit the induction of STAT3 phosphorylation by IFNα and IL-6 in breast cancer cells.		Floxuridine (5-Fluorouracil 2'-deoxyriboside) is a pyrimidine analog and known as an oncology antimetabolite .	of H o
Purity:99.78%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	.00 mg	Purity: 99.76% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 200 mg, 500 mg	HÔ
FLT3-IN-14	Cat. No.: HY-144777	FLT3/TrKA-IN-1	Cat. No. : HY-146749
FLT3-IN-14 is a potent FLT3 inhibitor with IC ₅₀ s of 5.6 nM and 1.4 nM for FLT3-WT and FLT3-ITD. FLT3-IN-14 reduces the phosphorylation of FLT3 (Y591), induces cell cycle arrest at G1 phase and apoptosis . FLT3-IN-14 significantly reduces the tumor growth is an MV4-11 papograft moure model	+0440-54	FLT3/TrKA-IN-1 is a potent FLT3/TrKA dual kinase inhibitor with the IC_{so} s of 43.8 nM, 97.2 nM, 92.5 nM and 23.6 nM for FLT3, FLT3-ITD, FLT3-TKD and TrKA, respectively.	
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	~N~~0~~
Flubendazole	Cat. No.: HY-B0294	Flubendazole-d3	Cat. No.: HY-B0294S
Flubendazole is a safe and efficacious anthelmintic drug, which is widely used for anthelmintic to human, rodents and ruminants. Flubendazole exerts anticancer activities by mechanisms including inhibition of microtubule function.	FLITCH N OF	Flubendazole-d3 is the deuterium labeled Flubendazole. Flubendazole is a safe and efficacious anthelmintic drug, which is widely used for anthelmintic to human, rodents and ruminants.	
Purity:99.79%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 100 mg, 500 mg		Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	

Fludarabine		Fludarabine phosphate	
(F-ara-A; NSC 118218)	Cat. No.: HY-B0069	(NSC 118218 phosphate)	Cat. No.: HY-B0028
Fludarabine (NSC 118218) is a DNA synthesis inhibitor and a fluorinated purine analogue with antineoplastic activity in lymphoproliferative malignancies.	H ₂ N N N OH F	Fludarabine (phosphate) is an analogue of adenosine and deoxyadenosine, which is able to compete with dATP for incorporation into DNA and inhibit DNA synthesis.	HO CH N N N HO C N N N N
Purity: 99.85% Clinical Data: Launched Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg	но	Purity: ≥98.0% Clinical Data: Launched Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg	0 он
Fludarabine triphosphate (F-ara-ATP)	Cat. No.: HY-136650	Fluorizoline	Cat. No.: HY-114989
Fludarabine triphosphate (F-ara-ATP), the cytotoxic metabolite of Fludarabine phosphate (HY-80028), inhibits ribonucleotide reductase and DNA polymerase and ultimately leads to cellular apoptosis.		Fluorizoline selectively and directly binds to prohibitin 1 (PHB1) and 2 (PHB2), and induces apoptosis . Fluorizoline reduces chronic lymphocytic leukemia (CLL) cell viability through the upregulation of NOXA and BIM. Fluorizoline exerts antitumor action in a p53-independent manner.	
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity: 99.77% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	00 mg
Flurbiprofen		Flurbiprofen-13C,d3	Cat No. UV 1059252
Flurbiprofen (dl-Flurbiprofen) is a potent, orally active nonsteroidal anti-inflammatory agent (NSAIA/NSAID), with antipyretic and analgesic activities.		Flurbiprofen-13C,d3 is the 13C- and deuterium labeled. Flurbiprofen (dl-Flurbiprofen) is a potent, orally active nonsteroidal anti-inflammatory agent (NSAIA/NSAID), with antipyretic and analgesic activities.	
Purity: 99.92% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 500 mg		Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	
Flurbiprofen-d3		Flurbiprofen-d5	
(dl-Flurbiprofen-d3)	Cat. No.: HY-10582S	(dl-Flurbiprofen-d5)	Cat. No.: HY-10582S1
Flurbiprofen-d3 (dl-Flurbiprofen-d3) is the deuterium labeled Flurbiprofen. Flurbiprofen (dl-Flurbiprofen) is a potent, orally active nonsteroidal anti-inflammatory agent (NSAIA/NSAID), with antipyretic and analgesic activities. Purity: >98% Clinical Data: No Development Reported Size: 5 mg, 50 mg	D D H H F F	Flurbiprofen-d5 (dl-Flurbiprofen-d5) is the deuterium labeled Flurbiprofen. Flurbiprofen (dl-Flurbiprofen) is a potent, orally active nonsteroidal anti-inflammatory agent (NSAIA/NSAID), with antipyretic and analgesic activities. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg, 50 mg	
Formononetin (Biochanin B: Flavosil: Formononetol)	Cat No : HY-N0183	Formosanin C	Cat No. HV-N2389
Formononetin is a potent FGFR2 inhibitor with an IC_{50} of ~4.31 μ M. Formononetin potently inhibits angiogenesis and tumor growth.	Ha	Formosanin C is a diosgenin saponin isolated from Paris formosana Hayata and an immunomodulator with antitumor activity. Formosanin C induces apoptosis.	
Purity: 99.88% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg	j, 500 mg	Purity:99.28%Clinical Data:No Development ReportedSize:1 mg, 5 mg, 10 mg, 25 mg	





Galanthamine		Galanthamine-d6	
(Galantamine)	Cat. No.: HY-76299		Cat. No.: HY-76299S
Galanthamine is a potent acetylcholinesterase (AChE) inhibitor with an $\mathrm{IC}_{\mathrm{so}}$ of 500 nM.	, where the second seco	Galanthamine-d6 (Galantamine-d6) is the deuterium labeled Galanthamine. Galanthamine is a potent acetylcholinesterase (AChE) inhibitor with an $\rm IC_{50}$ of 500 nM.	N N N N N N N N N N N N N N N N N N N
Purity: 99.90% Clinical Data: Launched Size: 10 mM × 1 mL, 50 mg, 100 mg	HO. ~ 0 °	Purity:>98%Clinical Data:Size:1 mg, 10 mg	HO. A HO A D
Galanthamine-O-methyl-d3	Cat. No.: HY-76299S1	Galgravin	Cat. No.: HY-N5007
Galanthamine-O-methyl-d3 is the deuterium labeled Galanthamine. Galanthamine is a potent acetylcholinesterase (AChE) inhibitor with an IC_{s0} of 500 nM.	HOLOGO	Galgravin is an active compound in Nectandra megapotamica, with anti-inflammatory activity. Galgravin displays in vitro cytotoxic activity and induce apoptosis in leukemia cells.	p-g-g
Purity:>98%Clinical Data:No Development ReportedSize:2.5 mg, 25 mg	n U	Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg	
Collinerid		Callia asid budrata	
Gallic acid (3,4,5-Trihydroxybenzoic acid)	Cat. No.: HY-N0523	Gallic acid nydrate (3,4,5-Trihydroxybenzoic acid hydrate)	Cat. No.: HY-N0523A
Gallic acid (3,4,5-Trihydroxybenzoic acid) is a natural polyhydroxyphenolic compound and an free radical scavenger to inhibit cyclooxygenase-2 (COX-2). Gallic acid has various activities, such as antimicrobial, antioxidant, antimicrobial, anti-inflammatory, and anticance activities.	но он	Gallic acid (3,4,5-Trihydroxybenzoic acid) hydrate is a natural polyhydroxyphenolic compound and an free radical scavenger to inhibit cyclooxygenase-2 (COX-2).	но он но он но он
Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 500 mg		Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg	1120
Ganetespib		Ganoderenic acid D	
(STA-9090)	Cat. No.: HY-15205		Cat. No.: HY-N1516
Ganetespib (STA-9090) is a heat shock protein 90 (HSP90) inhibitor which exhibits potent cytotoxicity in a wide variety of hematological and solid tumor cell lines. Ganetespib has antiangiogenic effects in colorectal cancer mediated through inhibition of HIF-1 α and STAT3.	HN N OH	Ganoderenic acid D is a triterpene identified from the effective compounds of Ganoderma lucidum extract (GLE). Ganoderenic acid D inhibits the proliferation of cancer cells by inducing cell cycle arrest and apoptosis .	сулсан Сараба
Purity: 99.84%	↓ OH	Purity: ≥99.0%	0 / · 0
Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg,	200 mg	Size: 1 mg, 5 mg	
Consideria anid A		Consideria asid D	
Ganoderic acid A	Cat. No.: HY-N1447	Ganoderic acid D	Cat. No.: HY-N1511
Ganoderic acid A can inhibit of the JAK-STAT3 signaling pathway, also inhibit proliferation, viability, ROS.	о с ц ц с н с о н с н с о н	Ganoderic acid D, a highly oxygenated tetracyclic triterpenoid, is the major active component of Ganoderma lucidum. Ganoderic acid D upregulates the protein expression of SIRT3 and induces the deacetylated cyclophilin D (CypD) by SIRT3.	оста ста оста оста оста оста оста оста о
Purity:99.84%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 25 mg	Landon and Anna and A	Purity:99.40%Clinical Data:No Development ReportedSize:1 mg, 5 mg	7.99

Consideric acid DM		Carrinal	
	Cat No: HY-120140	Garcinor	Cat No : HY-107569
Ganoderic acid DM, a natural triterpenoid isolated from Ganoderma lucidum, induces DNA damage, G1 cell cycle arrest and apoptosis in human breast cancer cells. Ganoderic acid DM as a specific inhibitor of osteoclastogenesis. Purity: 99.65% Clinical Data: No Development Reported Size: 1 mg, 5 mg	$ \begin{array}{c} \begin{array}{c} \begin{array}{c} \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\$	Garcinol, a polyisoprenylated benzophenone harvested from Garcinia indica, exerts anti-cholinesterase properties towards acetyl cholinesterase (AChE) and butyrylcholinesterase (BChE) with IC $_{50}$ s of 0.66 μ M and 7.39 μ M, respectively.Purity:98.85% Clinical Data:No Development Reported Size:10 mM × 1 mL, 1 mg, 5 mg	
Gardenin B	Cat. No.: HY-N6037	GDC-0623 (RG 7421; MEK inhibitor 1)	Cat. No.: HY-15610
Gardenin B is a flavonoid isolated from Baccharis scandens. Gardenin B induces cell death in human leukemia cells involves multiple caspases.		GDC-0623 (RG 7421) is a potent, ATP-uncompetitive inhibitor of MEK1 (K _i =0.13 nM, +ATP), and displays 6-fold weaker potency against HCT116 (KRAS (G13D), EC_{s0} =42 nM) versus A375 (BRAF ^{V600E} , EC_{s0} =7 nM).	HO. O HIN SHI
Purity:99.88%Clinical Data:No Development ReportedSize:5 mg, 10 mg		Purity: 99.15% Clinical Data: Phase 1 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
		CEN144	
GEIM-5	Cat. No.: HY-146540	GEM144	Cat. No.: HY-143411
GEM-5 is a gemcitabine-based conjugate containing a HIF-1 α inhibitor (YC-1) (IC ₅₀ =30 nM). GEM-5 can significantly down-regulate the expression of HIF-1 α and up-regulate the expression of tumor suppressor p53. GEM-5 induces the apoptosis of A2780 cells and inhibits tumor growth. Purity: >98% Clinical Data: No Development Reported	agerider	GEM144 is a potent and orally active DNA polymerase α (POLA1) and HDAC 11 dual inhibitor. GEM144 induces acetylation of p53, activation of p21, G1/S cell cycle arrest, and apoptosis. Purity: >98% Clinical Data: No Development Reported	но- но- но- но- но- с- с- с- с- с- с- с- с- с- с- с- с- с-
Size: 1 mg, 5 mg		Size: 1 mg, 5 mg	
Compitabing		Compitabino elaidata	
(LY 188011)	Cat. No.: HY-17026	(CP-4126; CO-101; Gemcitabine 5'-elaidate)	Cat. No.: HY-13538
Gemcitabine (LY 188011) is a pyrimidine nucleoside analog antimetabolite and an antineoplastic agent. Gemcitabine inhibits DNA synthesis and repair, resulting in autophagy and apoptosis .	F OH Han N O	Gemcitabine elaidate (CP-4126) is lipophilic pro-drug of Gemcitabine. Gemcitabine elaidate is converted to Gemcitabine by esterases in order to be phosphorylated. Gemcitabine elaidate exhibits anti-tumor activity.	
Purity: 99.92% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 200 mg, 500 mg, 1 g		Purity: 98.22% Clinical Data: Launched Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
Gemcitabine elaidate hydrochloride (CP-4126 hydr CO-101 hydrochloride;)	rochloride; Cat. No.: HY-13538A	Gemcitabine hydrochloride (LY 188011 hydrochloride)	Cat. No.: HY-B0003
Gemcitabine elaidate (CP-4126) hydrochloride is lipophilic pro-drug of Gemcitabine. Gemcitabine elaidate hydrochloride is converted to Gemcitabine by esterases in order to be phosphorylated. Gemcitabine elaidate hydrochloride exhibits anti-tumor activity.		Gemcitabine Hydrochloride (LY 188011 Hydrochloride) is a pyrimidine nucleoside analog antimetabolite and an antineoplastic agent. Gemcitabine Hydrochloride inhibits DNA synthesis and repair, resulting in autophagy and apoptosis .	
Purity: 297.0% Clinical Data: No Development Reported Size: 5 mg, 10 mg, 50 mg, 100 mg		Funct. 99.93% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 200 mg, 500 mg, 1 g	

Geniposidic acid	Cat. No.: HY-N0010	Genistein (NPI 031L)	Cat. No.: HY-14596
$ \begin{array}{ll} \mbox{Geniposidic acid is an effective anticancer and} \\ \mbox{radioprotection agent. Target: Others Mice were} \\ \mbox{given an intraperitoneal injection of Geniposidic} \\ \mbox{acid (GA) (12.5, 25, 50 mg/kg) 1 h before} \\ \mbox{receiving GA against d-galactosamine (GalN) (800 mg/kg)/LPS (40 \mug/kg). \\ \mbox{Purity:} & \geq 98.0\% \\ \mbox{Clinical Data:} No Development Reported \\ \mbox{Size:} & 10 mM \times 1 mL, 5 mg, 10 mg, 50 mg, 100 mg \\ \end{array} $		Genistein, a soy isoflavone, is a multiple tyrosine kinases (e.g., EGFR) inhibitor which acts as a chemotherapeutic agent against different types of cancer, mainly by altering apoptosis, the cell cycle, and angiogenesis and inhibiting metastasis. Purity: 99.84% Clinical Data: Phase 4 Size: 10 mM × 1 mL, 100 mg, 500 mg	
Genistein 8-c-glucoside (G8CG)	Cat. No.: HY-N6882	Genistein-d4 (NPI 031L-d4)	Cat. No.: HY-14596S
Genistein 8-c-glucoside (G8CG) is a glucoside. Genistein 8-c-glucoside induces mitochondrial membrane depolarization and induces apoptosis .		Genistein-d4 (NPI 031L-d4) is the deuterium labeled Genistein. Genistein, a soy isoflavone, is a multiple tyrosine kinases (e.g.	HO CH O D OH
Purity:99.40%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 1 mg, 5 mg	он о 🖉 он	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
Genistin (Genistine; Genistoside; Genistein 7-Ο-β-D-glucopyranoside)	Cat. No.: HY-N0595	Geranyl acetate	Cat. No.: HY-N7070
Genistin (Genistine), an isoflavone belonging to the phytoestrogen family, is a potent anti-adipogenic and anti-lipogenic agent. Genistin attenuates cellular growth and promotes apoptotic cell death breast cancer cells through modulation of ERalpha signaling pathway.	HOL PH A CON	Geranyl acetate, an acyclic monoterpene ester derived from geraniol, is widely used in the cosmetics industry due to its pleasant scent. Geranyl acetate can induces cell apoptosis .	lond
Purity: 98.04% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 20	10 mg	Purity:99.77%Clinical Data:No Development ReportedSize:1 g, 5 g	
GGTI-2154	Cat. No. : HY-16229	GGTI-2154 hydrochloride	Cat. No. : HY-16229A
GGTI-2154 is a potent and selective inhibitor of geranylgeranyltransferase I (GGTase I), with an IC ₅₀ of 21 nM. GGTI-2154 shows more than 200-fold selectivity for GGTase I over FTase (IC50=5600 nM). GGTI-2154 can be used for the research of cancer. Purity: >98% Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg	N NH H	GGTI-2154 hydrochloride is a potent and selective inhibitor geranylgeranyltransferase I (GGTase I), with an IC ₅₀ of 21 nM. GGTI-2154 hydrochloride shows more than 200-fold selectivity for GGTase I over FTase (IC50=5600 nM). GGTI-2154 hydrochloride can be used for the research of cancer. Purity: 98.13% Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
GGTI-2418	Cat. No .: HY-16231	GGTI298	Cat. No. : HY-100876
GGTI-2418 is a highly potent, competitive, and selective geranylgeranyltransferase I (GGTase I) inhibitor. GGTI-2418 inhibits GGTase I and FTase activities with IC _{so} s of 9.5 nM and 53 μ M, respectively.		GGTI298 is a CAAZ peptidomimetic geranylgeranyltransferase I (GGTase I) inhibitor, strongly inhibiting the processing of geranylgeranylated Rap1A with little effect on processing of farnesylated Ha-Ras, with IC ₅₀ values of 3 and > 20 μ M in vivo, respectively.	
Purity: 98.04% Clinical Data: Phase 1 Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100) mg	Purity: ≥95.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 25 mg, 50	mg

GGTI298 Trifluoroacetate	Cat. No. : HY-15871	Ginkgetin	Cat. No.: HY-N0889
GGTI298 Trifluoroacetate is a CAAZ peptidomimetic geranylgeranyltransferase I (GGTase I) inhibitor, which can inhibit Rap1A with IC ₅₀ of 3 μ M; little effect on Ha-Ras with IC ₅₀ of >20 μ M. Purity: \geq 98.0% Clinical Data: No Development Reported	на уда Старования На уда уда Старования На уда Старования На уда уда Старования На уда На уда Старования На уда уда Старования На уда уда	Ginkgetin, a biflavone, is isolated from Ginkgo biloba leaves. Ginkgetin exhibit anti-tumor, anti-inflammatory, neuroprotective, anti-fungal activities. Ginkgetin is also a potent inhibitor of Wnt signaling , with an IC _{so} of 5.92 μM. Purity: 99.53%	Contraction of the second seco
Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 25 mg, 50 mg	g	Size: 10 mM × 1 mL, 5 mg, 10 mg	
Ginkgolide B (BN-52021)	Cat. No.: HY-N0784	Ginsenoside F2	Cat. No. : HY-125848
Ginkgolide B (BN-52021), an important active terpenoid from Ginkgo biloba leaves, is reported to increase cell viability and decrease cell apoptosis.		Ginsenoside F2, a metabolite from Ginsenoside Rb1, induces apoptosis accompanied by protective autophagy in breast cancer stem cells.	
Purity: ≥98.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 50 mg		Purity:99.95%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 20 mg	
Ginsenoside F4	Cat. No.: HY-N2503	Ginsenoside F5	Cat. No.: HY-108277
Ginsenoside F4 (GF4), ginseng saponinis, isolated from notoginseng or red ginseng. Ginsenoside F4 (GF4) has inhibitory effect on human lymphocytoma JK cell by inducing its apoptosis .		Ginsenoside F5, from crude extracts of flower buds of Panax ginseng, remarkably inhibits the growth of HL-60 cells by the apoptosis pathway.	HO HH HI O HH
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	но у бу у он он он	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	HOYO
Ginsenoside Rg1 (Panaxoside A; Panaxoside Rg1)	Cat. No.: HY-N0045	Ginsenoside Rg6	Cat. No.: HY-N0907
Ginsenoside Rg1 is one of the major active components of ginseng. Ginsenoside Rg1 ameliorates the impaired cognitive function, displays promising effects by reducing cerebral A β levels. Ginsenoside Rg1 also reduces NF- κ B nuclear translocation.		Ginsenoside Rg6 inhibits TNF- α -induced NF-κB transcriptional activity with an IC ₅₀ of 29.34 μ M in HepG2 cells. Ginsenoside Rg6 also exhibits apoptosis -inducing effect.	
Purity: ≥ 98.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg	Сон	Purity:99.13%Clinical Data:No Development ReportedSize:1 mg, 5 mg	Сон
Ginsenoside Rh2 (20(S)-Ginsenoside Rh2; 20(S)-Rh2; Ginsenoside-Rh2)	Cat. No.: HY-N0605	Ginsenoside Rk1	Cat. No.: HY-N2515
Ginsenoside Rh2 induces the activation of caspase-8 and caspase-9 . Ginsenoside Rh2 induces cancer cell apoptosis in a multi-path manner.	Han of the second secon	Ginsenoside Rk1 is a unique component created by processing the ginseng plant (mainly Sung Ginseng, SG) at high temperatures. Ginsenoside Rk1 has anti-inflammatory effect, suppresses the activation of Jak2/Stat3 signaling pathway and NF-κB.	
Purity: ≥ 98.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg		Purity:99.90%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 20 mg	

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Girinimbine		GKK1032B	
Girinimbin (Girinimbin) is a carbazole alkaloid with a variety of biological effects. Girinimbine can induce apoptosis , and has antitrypanosomal, antiplatelet activity, antibacterial activity, anti-inflammatory, antioxidant and antitumor activities.		GKK1032B is an alkaloid compound that can be found in endophytic fungus Penicillium sp. GKK1032B can induce the apoptosis of human osteosarcoma MG63 cells through caspase pathway activation.	
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	ΎΥΙ ΉΪ
Glaucocalyxin A	Cat. No.: HY-N2112	Gliotoxin (Aspergillin)	Cat. No.: HY-N6727
Glaucocalyxin A, an ent-kauranoid diterpene from Rabdosia japonica var., induces apoptosis in osteosarcoma by inhibiting nuclear translocation of Five-zinc finger Glis 1 (GLI1) via regulating PI3K/Akt signaling pathway. Glaucocalyxin A has antitumor effect.	O H OH	Gliotoxin is a secondary metabolite, the most abundant mycotoxin secreted by A. fumigatus, inhibits the phagocytosis of macrophages and the immune functions of other immune cells .	PH O H N SSN O
Purity:99.38%Clinical Data:No Development ReportedSize:5 mg, 10 mg		Purity: 99.51% Clinical Data: No Development Reported Size: 5 mg	
GLP-2(rat)	Cat. No.: HY-P1142	Glucagon-Like Peptide (GLP) II, human	Cat. No.: HY-P1841
GLP-2(rat) is an intestinal growth factor. GLP-2(rat) stimulates cell proliferation and inhibits apoptosis. GLP-2(rat) enhances mucosal mass and function in residual small intestine after massive small bowel resection (MSBR).	HKOGSPECEMENLENLATHOPHNALIGTKITD	Glucagon-Like Peptide (GLP) II, human is a 33-amino acid peptide derived from the C-terminal of proglucagon and mainly produced by the intestinal L cells. Glucagon-Like Peptide (GLP) II, human stimulates intestinal mucosal growth and decreases apoptosis of enterocytes .	HADOSPEDEMENTLENLAAMEPHAAJETKITE
Purity: >98% Clinical Data: Phase 3 Size: 1 mg, 5 mg		Purity: >98% Clinical Data: No Development Reported Size: 5 mg, 10 mg	
GLUT4-IN-2	Cat. No. : HY-146980	Glychionide A	Cat. No.: HY-N8034
GLUT4-IN-2 is a potent and selective GLUT4 inhibitor with IC_{50} s of 11.4 μ M and 6.8 μ M for GLUT1 and GLUT4, respectively. GLUT4-IN-2 induces cell apoptosis and cell cycle arrest at GO/G1phase. GLUT4-IN-2 shows potent antitumor activity.	OLS NH	Glychionide A is a flavonoside that can be found in the roots of Glychirriza glabra. Glychionide A promotes apoptosis and autophagy of PANC-1 pancreatic cancer cells. Glychionide A can be used for the research of cancer.	HO OH OH OH OH
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg	
Glycitein (Glycetein)	Cat. No.: HY-N0016	Glycochenodeoxycholic acid (Chenodeoxycholylglycine)	Cat. No.: HY-N2334
Glycitein is a soybean (yellow cultivar) isoflavonoid; used in combination with other isoflavonoids such as genistein and daidzein to study apoptosis and anti-oxidation processes.	HO O O O O O O O O O O O	Glycochenodeoxycholic acid (Chenodeoxycholylglycine) is a bile acid formed in the liver from chenodeoxycholate and glycine. It acts as a detergent to solubilize fats for absorption and is itself absorbed.	
Purity:98.17%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg		Purity: ≥98.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg	

Glycochenodeovycholic acid sodium salt		Glycochenodeoxycholic acid-d4	
(Chenodeoxycholylglycine sodium salt;)	Cat. No.: HY-N2334A	(Chenodeoxycholylglycine-d4)	Cat. No.: HY-N2334S
Glycochenodeoxycholic acid sodium salt (Chenodeoxycholylglycine sodium salt) is a bile acid formed in the liver from chenodeoxycholate and glycine. It acts as a detergent to solubilize fats for absorption and is itself absorbed. Purity: ≥98.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 100 mg		Glycochenodeoxycholic acid-d4 (Chenodeoxycholylglycine-d4) is the deuterium labeled Glycochenodeoxycholic acid. Glycochenodeoxycholic acid (Chenodeoxycholylglycine) is a bile acid formed in the liver from chenodeoxycholate and glycine. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg	аф Ца ho b di b di b di b di
Glycochenodeoxycholic acid-d7 sodium (Chenodeoxycholylglycine-d7 sodium;)	Cat. No.: HY-N2334AS	Glyphosate	Cat. No.: HY-B0863
Glycochenodeoxycholic acid-d7 (Chenodeoxycholylglycine-d7) sodium is the deuterium labeled Glycochenodeoxycholic acid (sodium salt).		Glyphosate is an herbicidal derivative of the amino acid glycine. Glyphosate targets and blocks a plant metabolic pathway not found in animals, the shikimate pathway, required for the synthesis of aromatic amino acids in plants.	о но он
Purity:> 98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity: ≥98.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 500 mg	
Glyphosate-d2		GMB-475	
	Cat. No.: HY-B0863S		Cat. No.: HY-125834
Glyphosate-d2 is the deuterium labeled Glyphosate. Glyphosate is an herbicidal derivative of the amino acid glycine. Glyphosate targets and blocks a plant metabolic pathway not found in animals, the shikimate pathway, required for the synthesis of aromatic amino acids in plants.		GMB-475 is a degrader of BCR-ABL1 tyrosine kinase based on PROTAC, overcoming BCR-ABL1-dependent drug resistance. GMB-475 targets BCR-ABL1 protein and recruits the E3 ligase Von Hippel Lindau (VHL), resulting in ubiquitination and subsequent degradation of the oncogenic fusion protein.	torante
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Purity:99.20%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg	
Gomisin N		Goserelin	
	Cat. No.: HY-N6866	(ICI 118630)	Cat. No.: HY-13673
Gomisin N, isolated from Schisandra chinensis. Gomisin N has the potential for use in the treatment of allergy. Gomisin N is an anti-cancer drug candidate capable of inhibiting the proliferation and inducing the apoptosis in cancer.		Goserelin (ICI 118630), a decapeptide analogue of gonadotropin-releasing hormone (GnRH/LHRH), functions as a GnRH agonist. Goserelin can be used for the research of breast cancer, epithelial ovarian cancer and prostate cancer.	
Purity:99.64%Clinical Data:No Development ReportedSize:5 mg, 10 mg	' ó.	Purity:>98%Clinical Data:LaunchedSize:1 mg, 5 mg	
Gosoralin acetata		GDI GIAGO	
(ICI-118630 acetate)	Cat. No.: HY-13673A	Gr Lainad	Cat. No.: HY-P2213
Goserelin acetate (ICI-118630 acetate), a decapeptide analogue of gonadotropin-releasing hormone (GnRH/LHRH), functions as a GnRH agonist. Goserelin acetate can be used for the research of breast cancer, epithelial ovarian cancer and prostate cancer.	a the	GPLGIAGQ, a MMP2-cleavable polypeptide, is used as a stimulus-sensitive linker in both liposomal and micellar nanocarriers for MMP2-triggered tumor targeting. GPLGIAGQ can be used to synthesis unique MMP2-targeted photosensitizer in photodynamic therapy (PDT).	La titte a
Purity: 99.89% Clinical Data: Launched Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg	. J.,	Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	

GPLGIAGQ TFA	Cat. No.: HY-P2213A	GPNA hydrochloride	Cat. No.: HY-W011391
GPLGIAGQ TFA, a MMP2-cleavable polypeptide, isused as a stimulus-sensitive linker in bothliposomal and micellar nanocarriers forMMP2-triggered tumor targeting. GPLGIAGQ TFA canbe used to synthesis unique MMP2-targetedphotosensitizer in photodynamic therapy (PDT).Purity:99.67%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg	(1) (1) (1) (1) (1) (1) (1) (1) (1) (1)	GPNA hydrochloride is a well known substrate of the enzyme γ-glutamyltransferase (GGT). GPNA hydrochloride is a specific glutamine (GIn) transporter ASCT2 inhibitor. Purity: 99.91% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg	O N H-CI
Grape seed extract	Cat. No.: HY-N7072	Griseofulvin	Cat. No.: HY-17583
Grape seed extract is a natural product, with anti-inflammatory and anti-proliferative effects. Grape seed extract shows inhibitory activity on the fat-metabolizing enzymes pancreatic lipase and lipoprotein lipase. Grape seed extract induces apoptotic in human colorectal cancer cells.	Grape seed extract	Griseofulvin(Gris-PEG; Grifulvin) is a spirocyclic fungal natural product used in treatment of fungal dermatophytes; Antifungal drug.	
Purity:>98%Clinical Data:Phase 3Size:100 mg, 250 mg, 500 mg		Purity: 98.89% Clinical Data: Launched Size: 10 mM × 1 mL, 500 mg, 5 g	
Griseofulvin-13C d3		Griseofulvin-d3	
	Cat. No.: HY-17583S1		Cat. No.: HY-17583S
Griseofulvin-13C,d3 is the 13C- and deuterium labeled.	0=	Griseofulvin-d3 is the deuterium labeled Griseofulvin, Griseofulvin (Gris-PEG) is a spirocyclic fungal natural product used in treatment of fungal dermatophytes; Antifungal drug.	
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	-	Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	
GRP78-IN-1		GS-444217	
	Cat. No.: HY-145857		Cat. No.: HY-100844
GRP78-IN-1 exhibits several interactions with GRP78 residues with binding energy of -8.07 kcal/mol. GRP78-IN-1 shows the potent cytotoxic, anti-proliferative in cancer cells. GRP78-IN-1 exhibits promising apoptosis in breast cancer cells and wound healing properties.		GS-444217 is a potent, orally available and selective ATP-competitive inhibitor of apoptosis signal-regulating kinase 1 (ASK1) with an IC _{s0} of 2.87 nM.	
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	0	Purity: 99.67% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	.00 mg
GSK 3 Inhibitor IX (6-Bromoindirubin-3'-oxime; BIO; MLS 2052)	Cat. No.: HY-10580	GSK-1070916 (GSK-1070916A)	Cat. No.: HY-70044
GSK 3 Inhibitor IX (6-Bromoindirubin-3'-oxime; BIO) is a potent, selective, reversible and ATP-competitive inhibitor of GSK-3 α/β and CDK1-cyclinB complex with IC ₅₀ s of 5 nM/320 nM/80 nM for (GSK-3 α/β)/CDK1/CDK5, respectively.		GSK-1070916 is a potent and selective ATP-competitive inhibitor of aurora B and aurora C with K _i s of 0.38 and 1.5 nM, respectively, and is >250- fold selective over Aurora A.	YE CHARGE
Purity: 99.74% Clinical Data: Phase 4 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg	0	Purity: 99.55% Clinical Data: Phase 1 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg,	200 mg





GZD856 formic **HA15** Cat. No.: HY-101489A Cat. No.: HY-100437 GZD856 formic is a potent and orally active HA15 is a potent and specific inhibitor of ER **PDGFR** α / β inhibitor, with IC₅₀s of 68.6 and 136.6 chaperone BiP/GRP78/HSPA5, inhibits the ATPase nM, respectively. GZD856 formic is also a activity of BiP, with anti-cancerous activity. Bcr-Abl^{T315I} inhibitor, with IC₅₀s of 19.9 and 15.4nM for native Bcr-Abl and the T315I mutant. GZD856 formic has antitumor activity. Purity: 98.06% 99 62% Purity: Clinical Data: No Development Reported Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg Haemanthamine Haemanthamine hydrochloride Cat. No.: HY-114489A Cat. No.: HY-114489B Haemanthamine is a crinine-type alkaloid isolated Haemanthamine hydrochloride is a crinine-type from the Amaryllidaceae plants with potent alkaloid isolated from the Amaryllidaceae anticancer activity. Haemanthamine targets plants with potent anticancer activity. ribosomal that inhibits protein biosynthesis Haemanthamine hydrochloride targets ribosomal that during the elongation stage of translation. inhibits protein biosynthesis during the elongation stage of translation. H-CI Purity: > 98% **Purity:** >98% Clinical Data: No Development Reported Clinical Data: No Development Reported 5 mg, 10 mg 5 mg, 10 mg, 25 mg Size: Size: HBDDE HBX 41108 Cat. No.: HY-131305 Cat. No.: HY-101666 HBDDE, a derivative of Ellagic acid, is an HBX 41108 is an uncompetitive inhibitor of isoform-selective PKCa and PKCy inhibitor with ubiquitin-specific protease 7 (USP7) with an IC₅₀ IC_{co}s of 43 μM and 50 μM, respectively. HBDDE of 424 nM. HBX 41108 inhibits USP7-mediated p53 shows selective for PKCα/PKCy over PKCδ, PKCβI deubiquitination to stabilize p53 and inhibits HC and PKCBII isozymes. HBDDE induces neuronal cancer cell growth. OH apoptosis. Ò. Purity: >98% >98% Purity: Clinical Data: No Development Reported Clinical Data: No Development Reported Size: 1 mg, 5 mg Size: 1 mg, 5 mg hCAIX/XII-IN-1 HDAC-IN-31 Cat. No.: HY-146988 Cat. No.: HY-144293 hCAIX/XII-IN-1 is a potent CAIX/XII inhibitor with HDAC-IN-31 is a potent, selective and orally the K, values of 0.48 μM and 0.83 μM for CAIX and active HDAC inhibitor with IC50 s of 84.90, 168.0, CAXII, respectively. hCAIX/XII-IN-1 shows 442.7, >10000 nM for HDAC1, HDAC2, HDAC3, antiproliferative activity in vitro. HDAC8, respectively. HDAC-IN-31 induces apoptosis hCAIX/XII-IN-1 induces apoptosis in MCF-7 cells. and cell cycle arrests at G2/M phase. HDAC-IN-31 shows good antitumor efficacy. Purity: >98% >98% Purity: Clinical Data: No Development Reported Clinical Data: No Development Reported Size: 1 ma, 5 ma Size 1 mg, 5 mg HDAC-IN-37 HDAC-IN-36 Cat. No.: HY-146684 Cat. No.: HY-146750 HDAC-IN-36 (compound 23 g) is an orally active and HDAC-IN-37 is a potent HDAC inhibitor with IC as potent HDAC (histone deacetylase) inhibitor, with of 0.0551 μM, 1.24 μM, 0.948 μM and 34.2 μM for an IC_{so} of 11.68 nM (HDAC6). HDAC-IN-36 promotes HDAC1, HDAC3, HDAC8 and HDAC6, respectively. apoptosis, autophagy and suppresses migration. HDAC-IN-37 induces histone acetylation in a slow-off manner. Purity: >98% Purity: >98% Clinical Data: No Development Reported Clinical Data: No Development Reported Size: 1 mg, 5 mg Size: 1 mg, 5 mg

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HDAC-IN-9		HDAC1/2 and CDK2-IN-1	
HDAC-IN-9 is a potent and selective tubulin and HDAC dual inhibitor. HDAC-IN-9 inhibits the invasion and migration of A549 cells. HDAC-IN-9 shows potent antitumor and antiangiogenic effect in vitro and in vivo. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	Cat. No.: HY-115941	$\label{eq:host} \begin{array}{l} \text{HDAC1/2 and CDK2-IN-1 (compound 14d) is a potent} \\ \text{HDAC1, HDAC2 and CDK2 dual inhibitor, with IC}_{50} \\ \text{values of 70.7, 23.1 and 0.80 } \mu\text{M}, \text{ respectively.} \\ \text{HDAC1/2 and CDK2-IN-1 can block the cell cycle and} \\ \text{induce apoptosis. HDAC1/2 and CDK2-IN-1 exhibits} \\ \text{desirable in vivo antitumor activity.} \\ \begin{array}{lllllllllllllllllllllllllllllllllll$	Cat. No.: HY-143497
HDAC6-IN-4	Cat No . HV. 14/305	HDACs/mTOR Inhibitor 1	Cat No : HV-114414
HDAC6-IN-4 (C10) is a potent, orally active and highly selective HDAC6 inhibitor with an IC ₅₀ value of 23 nM. HDAC6-IN-4 induces cancer cells apoptosis and shows significant antitumor efficacy, without obvious toxicity. Purity: >98% Clinical Data: No Development Reported	Office Cal. NO. HT-144395	HDACs/mTOR Inhibitor 1 is a dual Histone Deacetylases (HDACs) and mammalian target of Rapamycin (mTOR) target inhibitor for treating hematologic malignancies, with $IC_{so}s$ of 0.19 nM, 1.8 nM, 1.2 nM and >500 nM for HDAC1, HDAC6, mTOR and PI3K α , respectively. Purity: 98.21% Clinical Data: No Development Reported	Cat. NO.: HT-114414
Size: 1 mg, 5 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
Hederacolchiside A1	Cat. No. : HY-N6950	Helichrysetin	Cat. No. : HY-N4058
Hederacolchiside A1, isolated from Pulsatilla chinensis, suppresses proliferation of tumor cells by inducing apoptosis through modulating PI3K/Akt/mTOR signaling pathway.		Helichrysetin, isolated from the flowers of Helichrysum odoratissimum, is an ID2 (inhibitor of DNA binding 2) inhibitor, and suppresses DCIS (ductal carcinoma in situ) formation.	рн р но-С-о-О-он
Purity:99.69%Clinical Data:No Development ReportedSize:5 mg, 10 mg	NOTO	Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg	
Hellebrigenin	Cat. No.: HY-N6576	Hematein	Cat. No.: HY-119751
Hellebrigenin, one of bufadienolides belonging to cardioactive steroids, is isolated from traditional Chinese medicine Venenum Bufonis. Hellebrigenin induces DNA damage and cell cycle G2/M arrest. Hellebrigenin triggers mitochondria-mediated apoptosis. Purity: >98% Clinical Data: No Development Reported Size: 1 mg	HO OH	$\begin{array}{ll} \mbox{Hematein is a oxidation product of hematoxylin} \\ \mbox{acted as a dye. Hematein is an allosteric casein} \\ \mbox{kinase II inhibitor with an IC}_{50} of 0.74 \ \mu\text{M}. \\ \mbox{Hematein inhibits Akt/PKB Ser129} \\ \mbox{phosphorylation, the Wnt/TCF pathway and} \\ \mbox{increases apoptosis in lung cancer cells.} \\ \mbox{Purity:} & 74.90\% \\ \mbox{Clinical Data:} \\ \mbox{Size:} & 10 \ \text{mM} \times 1 \ \text{mL}, 500 \ \text{mg}, 1 \ \text{g} \end{array}$	но с с с с с с с с с с с с с с с с с с с
Hematoporphyrin (Hematoporphyrin IX) Hematoporphyrin (Hematoporphyrin IX), a	Cat. No.: HY-B0754	Hematoporphyrin dihydrochloride (Hematoporphyrin IX dihydrochloride) Hematoporphyrin dihydrochloride (Hematoporphyrin	Cat. No.: HY-B0754A
photosensitizer, is a substrate for affinity chromatography of heme-binding proteins. Hematoporphyrin can induce apoptosis in U87 glioma cells and decrease tumor growth in vivo when exposed to red light.		IX dihydrochloride), a photosensitizer, is a substrate for affinity chromatography of heme-binding proteins.	HO H-CI
Purity: 95.81% Clinical Data: No Development Reported Size: 100 mg		Purity:95.81%Clinical Data:Phase 1Size:100 mg	H-CI





HS-173		HS-1793	
	Cat. No.: HY-15868		Cat. No.: HY-129156
HS-173 is a novel PI3K inhibitor, that is used for cancer treatment.	Contraction of the	HS-1793 is a resveratrol analogue with antitumor activities in a variety of cancer cell lines. HS-1793 induces cell apoptosis .	носто
Purity:99.04%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
Hsp90-Cdc37-IN-3	Cat. No. : HY-144650	Humulone (α-Lupulic acid)	Cat. No. : HY-N6084
Hsp90-Cdc37-IN-3 (Compound 9) is a novel celastrol–imidazole derivative with anticancer activity. Hsp90-Cdc37-IN-3 inhibits H sp90-Cdc37 by covalent-binding, and induces apoptosis . Purity: >98% Clinical Data: No Development Reported	HO TO THE CONTRACT OF THE CONTRACT.	Humulone (α-Lupulic acid), a prenylated phloroglucinol derivative, is a potent cyclooxygenase-2 (COX-2) inhibitor. Humulone acts as a positive modulator of GABA, receptor at low micromolar concentrations. Humulone is an inhibitor of bone resorption. Purity: >98% Clinical Data: No Development Reported	HO CH
Size: 1 mg, 5 mg		Size: 5 mg, 10 mg, 25 mg	
HXR9	Cat. No.: HY-P3245	HXR9 hydrochloride	Cat. No.: HY-P3245A
HXR9 is a cell-permeable peptide and a competitive antagonist of HOX/PBX interaction. HXR9 antagonizes the interaction between HOX and a second transcrip-tion factor (PBX), which binds to HOX proteins in paralogue groups1 to 8. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	WYPWMKKHHRRRRRRR	HXR9 hydrochloride is a cell-permeable peptide and a competitive antagonist of HOX/PBX interaction.HXR9 hydrochloride antagonizes the interaction between HOX and a second transcrip-tion factor (PBX), which binds to HOX proteins in paralogue groups1 to 8.Purity:99.50% Clinical Data: No Development Reported Size:1 mg, 5 mg, 10 mg	WYPWMKCHHRRRRRRR (HCI sat)
Hydrolyzed Fumonisin B1 (Aminopentol)	Cat. No.: HY-N6730	Hydroxyurea (Hydroxycarbamide)	Cat. No.: HY-B0313
Hydrolyzed Fumonisin B1 (Aminopentol) is the backbone and main hydrolysis product of the mycotoxin Fumonisin B1 (HY-N6719). Hydrolyzed Fumonisin B1 can weakly inhibit ceramide synthase.	~~~~ ⁰¹ be 1 be 400	Hydroxyurea is a cell apoptosis inducer that inhibit DNA synthesis through inhibition of ribonucleotide reductase .	
Purity: >98% Clinical Data: No Development Reported Size: 1 mg		Purity: ≥ 98.0% Clinical Data: Launched Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g	
Hypericin	Cat. No.: HY-N0453	Hypericin-d10	Cat. No.: HY-N0453S
Hypericin is a photosensitive antiviral with anticancer and antidepressant agent derived from Hypericum perforatum. It can inhibit tyrosine kinases with IC50 of 7.5 μ M.	HO HO OH HO HO HO	Hypericin-d10 is the deuterium labeled Hypericin. Hypericin is a photosensitive antiviral with anticancer and antidepressant agent derived from Hypericum perforatum. It can inhibit tyrosine kinases with IC ₅₀ of 7.5 μ M.	
Purity: ≥98.0% Clinical Data: Phase 1 Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 25 mg	ү ү ү он о он	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	он о он

Hypericin-d2		Hypocrellin B	
	Cat. No.: HY-N0453S1		Cat. No.: HY-N1453
Hypericin-d2 is deuterium labeled Hypericin. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Hypocrellin B, a pigment isolated from the fungi Hypocrella bambusae and Shiraia bambusicola, is an apoptosis inducer. Hypocrellin B can be used as a photosensitizer for photodynamic therapy of cancer. Hypocrellin B also has antimicrobial and antileishmanial activities.Purity:99.61% Clinical Data:No Development Reported Size:5 mg, 10 mg	
IACS-010759 (IACS-10759)	Cat. No.: HY-112037	IACS-010759 hydrochloride (IACS-10759 hydrochloride)	Cat. No.: HY-112037A
IACS-010759 is an orally active, potent mitochondrial complex I of oxidative phosphorylation (OXPHOS) inhibitor. IACS-010759 inhibits proliferation and induces apoptosis in models of brain cancer and acute myeloid leukemia (AML) reliant on OXPHOS.	9074008	IACS-010759 hydrochlorideis an orally active, potent mitochondrial complex I of oxidative phosphorylation (OXPHOS) inhibitor.	No Contraction
Clinical Data: Phase 1		Clinical Data: Phase 1	
Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg,	100 mg	Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg,	100 mg
Ibandronate Sodium Monohydrate		Iberdomide	
(BM-210955; RPR-102289A)	Cat. No.: HY-B0515	(CC-220)	Cat. No.: HY-101291
Ibandronate Sodium Monohydrate is a highly potent nitrogen-containing bisphosphonate used for the treatment of osteoporosis.	, он Р. он 0::P. он NaO H ₂ O	Iberdomide (CC-220) is an orally active and potent cereblon (CRBN) E3 ligase modulator (CELMoD) with an IC ₅₀ of ~150nM for cereblon-binding affinity. Iberdomide, a derivative of Thalidomide (HY-14658), has antitumor and immunostimulatory activities.	S S S S S S S S S S S S S S S S S S S
Clinical Data: Launched		Clinical Data: Phase 3	•
Size: 10 mM × 1 mL, 100 mg, 500 mg		Size: 10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 25 mg, 5	0 mg, 100 mg
Iberin		IBR2	
(NSC 321801)	Cat. No.: HY-101413		Cat. No.: HY-103710
Iberin (NSC 321801), a sulfoxide analogue of sulforaphane, is a naturally occurring member of isothiocyanate family. Iberin inhibits cell survival with an IC_{s0} of 2.3 μ M in HL60 cell. Iberin induces apoptosis . Purity: 98.0% Clinical Data: No Development Reported Size: 1 mg (61.25 mM * 100 μ L in Ethanol),	S ^{2C²N}	IBR2 is a potent and specific RAD51 inhibitor and inhibits RAD51-mediated DNA double-strand break repair. IBR2 disrupts RAD51 multimerization, accelerates proteasome-mediated RAD51 protein degradation, inhibits cancer cell growth and induces apoptosis. Purity: 98.60% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	NH N ^S O
IC261	Cat. No. : HY-12774	Icariside D2	Cat. No.: HY-N7450
IC261 is a selective, ATP-competitive CK1 inhibitor, with IC ₅₀ S of 1 μ M, 1 μ M, 16 μ M for Ckiδ, Ckiε and Ckiα1, respectively.	H P P P P	Icariside D2, isolated from Annona glabra fruit, inhibits angiotensin-converting enzyme . Icariside D2 shows significant cytotoxic activity on the HL-60 cell line with the IC ₅₀ value of 9.0 \pm 1.0 μ M. Icariside D2 induces apoptosis .	но обо он он
Purity:99.75%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg	-0-0	Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg	





Indirubin		Inecalcitol	
(Couroupitine B; Indigo red; Indigopurpurin)	Cat. No.: HY-N0117	(TX 522)	Cat. No.: HY-32344
Indirubin (Couroupitine B) is a purple 3,2- bisindole and a stable isomer of indigo isolated from Indigo naturalis (Apiaceae); anti-inflammatory and anticancer activities.	HN CO	Inecalcitol (TX 522), a unique vitamin D3 analog, is an orally active vitamin D receptor (VDR) agonist with a K_a of 0.53 nM. Inecalcitol can induce cell apoptosis and has potent anticancer activities.	ноу то в стран
Purity: ≥98.0% Clinical Data: Launched Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100) mg, 250 mg	Purity:98.11%Clinical Data:No Development ReportedSize:1 mg, 5 mg, 10 mg	
Infigratinib (BGJ-398; NVP-BGJ398)	Cat. No .: HY-13311	Ingenol 3,20-dibenzoate	Cat. No.: HY-137295
Infigratinib (BGJ-398; NVP-BGJ398) is a potent inhibitor of the FGFR family with IC ₅₀ s of 0.9 nM, 1.4 nM, 1 nM, and 60 nM for FGFR1, FGFR2, FGFR3, and FGFR4, respectively.	<i>oobab</i> ę	Ingenol 3,20-dibenzoate is a potent protein kinase C (PKC) isoform-selective agonist.	
Purity: 99.70% Clinical Data: Launched Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 20)0 mg	Purity:99.31%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg	0
INH1		INH6	
(IB113131)	Cat. No.: HY-16660		Cat. No.: HY-100541
INH1 specifically disrupts the Hec1/Nek2 interaction via direct Hec1 binding. INH1 shows promising cancer inhibition activity both in vitro and in vivo.	CC NH O	INH6 is a potent Nek2/Hec1 inhibitor; inhibits the growth of HeLa cells with an IC $_{\rm s0}$ of 2.4 $\mu M.$	LL N-NH
Purity: 99.62% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 500 mg		Purity:99.38%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg, 25 mg, 50 mg, 100 mg	
lodoacetyl-LC-biotin	Cat No: HY-138065	lonomycin (SO23377)	Cat No : HY-13434
Iodoacetyl-LC-biotin is a biotinylated electrophile probe that can be used to investigate the scope and characteristics of protein covalent binding to subcellular proteomes.	,	Ionomycin (SQ23377) is a potent, selective calcium ionophore and an antibiotic produced by Streptomyces conglobatus. Ionomycin (SQ23377) is highly specific for divalent cations (Ca>Mg>Sr=Ba). Ionomycin (SQ23377) promotes apoptosis. Purity: >99.0%	Martin time
Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg		Clinical Data: No Development Reported Size: 10 mg (14.1 mM * 1 mL in Ethanol)	
Ionomycin calcium (SQ23377 calcium)	Cat. No.: HY-13434A	Irbesartan (SR-47436; BMS-186295)	Cat. No.: HY-B0202
Ionomycin calcium (SQ23377 calcium) is a potent, selective calcium ionophore and an antibiotic produced by Streptomyces conglobatus. Ionomycin calcium (SQ23377 calcium) is highly specific for divalent cations (Ca>Mg>Sr=Ba). Ionomycin (SQ23377) promotes apoptosis .	magnathant	Irbesartan is a highly potent and specific angiotensin II type 1 (AT1) receptor antagonist with IC50 of 1.3 nM.	Grad And And And And And And And And And An
Purity:98.0%Clinical Data:No Development ReportedSize:1 mg, 5 mg, 10 mg		Purity: 98.98% Clinical Data: Launched Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg,	500 mg

Irbesartan-d6		Isatin	Cot No - UV VODCE
Irbesartan-d6 is the deuterium labeled Irbesartan. Irbesartan is a highly potent and specific angiotensin II type 1 (AT1) receptor antagonist with IC_{50} of 1.3 nM.		Isatin (Indoline-2,3-dione) is a potent inhibitor of monoamine oxidase (MAO) with an IC ₅₀ of 3 μM. Also binds to central benzodiazepine receptors (IC ₅₀ against clonazepam, 123 μM).	
Clinical Data: No Development Reported Size: 1 mg, 5 mg	HN-N	Clinical Data: No Development Reported Size: 10 mM × 1 mL, 100 mg	
Isatuximab	Cat. No.: HY-P9976	Isoalantolactone ((+)-Isoalantolactone; Isohelenin)	Cat. No.: HY-N0780
Isatuximab is a monoclonal antibody targeting the transmembrane receptor and ectoenzyme CD38 , a protein highly expressed on hematological malignant cells, including those in multiple myeloma (MM).	Isatuximab	Isoalantolactone is an apoptosis inducer, which also acts as an alkylating agent.	
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg, 25 mg, 50 mg		Purity:99.99%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg, 50 mg	
Isobavachalcone (Corylifolinin; Isobacachalcone)	Cat. No. : HY-13065	Isocurcumenol	Cat. No.: HY-N4121
Isobavachalcone (Corylifolinin) is derived from Psoralea corylifolia Linn. and is a potent inhibitor of Akt signaling pathway, which induces apoptosis in human cancer cells (Inhibits OVCAR-8 cell growth with an IC _{s0} value of 7.92 μ M).	HO-CH B	Isocurcumenol, an estrogen receptor alpha (ERα) inhibitor isolated from Curcuma zedoaria Rhizomes, possesses anti-tumor acticity, with IC ₅₀ values of 99.1µg/mL and 178.2 µg/mL in DLA and KB cells, respectively.	но
Purity:99.01%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 25 mg		Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg	// н
Isofistularin-3	Cat. No .: HY-19826	Isoliensinine	Cat. No.: HY-N0770
Isofistularin-3 is a direct, DNA-competitive DNMT1 inhibitor, with an IC _{s0} of 13.5 µM. Isofistularin-3, as a DNA demethylating agent, induces cell cycle arrest and sensitization to TRAIL in cancer cells. Isofistularin-3 can be used as an ADC cytotoxin. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	and the and th	Isoliensinine is a bisbenzylisoquinoline alkaloid extracted from the seed embryo of Nelumbo nucifera, with anti-oxidant and anti-inflammatory and anti-cancer activities. Isoliensinine induces apoptosis in triple-negative human breast cancer cells.Purity:99.83%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 20 mg	HOLL N OL
Isoliquiritigenin (GU17; ISL; Isoliquiritigen)	Cat. No.: HY-N0102	Isolongifolene ((-)-Isolongifolene)	Cat. No.: HY-N7363
Isoliquiritigenin is an anti-tumor flavonoid from the root of Glycyrrhiza glabra, which inhibits aldose reductase with an IC ₅₀ of 320 nM. Isoliquiritigenin is a potent inhibitor of influenza virus replication with an EC ₅₀ of 24.7 μ M.	ностон	Isolongifolene ((-)-Isolongifolene) is a tricyclic sesquiterpene isolated from Murraya koenigii. Isolongifolene attenuates Rotenone-induced oxidative stress, mitochondrial dysfunction and apoptosis through the regulation of PI3K/AKT/GSK-3β signaling pathways.	H
Purity: 98.17% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg	9	Purity: ≥98.0% Clinical Data: No Development Reported Size: 5 mg, 10 mg	<i>i</i> N

Isosilybin A		Isosilybin B	
Isosilybin A, a flavonolignan isolated from silymarin, has anti-prostate cancer (PCA) activity.	Cat. No.: HY-N/043	Isosilybin B, a flavonolignan isolated from silymarin, has anti-prostate cancer (PCA) activity via inhibiting proliferation and inducing G1 phase arrest and apoptosis . Isosilybin B causes androgen receptor (AR) degradation.	Cat. No.: HY-N/045
Purity: ≥99.0% Clinical Data:	900-900	Purity: 99.32% Clinical Data:	1972 - ANG
Ispinesib (SB-715992)	Cat. No.: HY-50759	ISRIB (trans-isomer)	Cat. No.: HY-12495
Ispinesib is a specific inhibitor of kinesin spindle protein (KSP), with a $K_{i app}$ of 1.7 nM.		ISRIB (trans-isomer) is a potent inhibitor of PERK with an IC ₅₀ of 5 nM. ISRIB potently reverses the effects of eIF2 α phosphorylation (IC ₅₀ =5 nM).	and the sense sense terrary
Purity: 99.74% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity:99.37%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg, 50 mg, 100 mg	
IV-23	Cat. No.: HY-126324	Ivaltinostat (CG-200745)	Cat. No.: HY-16138
IV-23 (Compound 20) is a potent Noxa mediated apoptosis inducer, and it is a promising anticancer agent with potential. IV-23 inhibits cell growths in vitro and in vivo, reduces colony formation, arrests cell cycle at M phase, and induces esophageal squamous cell carcinoma (ESCC). Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	Br O O O O O O O O O O O O O O O O O O O	Ivaltinostat (CG-200745) is an orally active, potent pan-HDAC inhibitor which has the hydroxamic acid moiety to bind zinc at the bottom of catalytic pocket. Ivaltinostat inhibits deacetylation of histone H3 and tubulin. Purity: >98% Clinical Data: Phase 2 Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
Ivaltinostat formic (CG-200745 formic)	Cat. No.: HY-16138A	IVHD-valtrate	Cat. No.: HY-N3446
Ivaltinostat (CG-200745) formic is an orally active, potent pan- HDAC inhibitor which has the hydroxamic acid moiety to bind zinc at the bottom of catalytic pocket. Ivaltinostat formic inhibits deacetylation of histone H3 and tubulin.	ц , , , , , , , , , , , , ,	IVHD-valtrate, an active Valeriana jatamansi derivative, is against human ovarian cancer cells in vitro and in vivo. IVHD-valtrate induces cancer cells apoptosis and arrests the ovarian cancer cells in the G2/M phase.	
Purity:99.36%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg		Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg	*
Ixabepilone (BMS-247550; Aza-epothilone B)	Cat. No.: HY-10222	Jaceosidin	Cat. No.: HY-N0831
Ixabepilone (BMS-247550) is an orally bioavailable microtubule inhibitor, which binds to tubulin and promotes tubulin polymerization and microtubule stabilization, thereby arrests cells in the G2-M phase of the cell cycle and induces tumor cell apoptosis.	N HN HOL	Jaceosidin is a flavonoid isolated from Artemisia vestita, induces apoptosis in cancer cells, activates Bax and down-regulates Mcl-1 and c-FLIP expression.	HO OH O
Purity: 99.93% Clinical Data: Launched Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg		Purity:99.51%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 25 mg	





Kaempferol Kahweol (Kempferol; Robigenin) Cat. No.: HY-14590 Cat. No.: HY-N6258 Kaempferol (Kempferol), a flavonoid found in many Kahweol is one of the consituents of the coffee edible plants, inhibits estrogen receptor α from Coffea Arabica with anti-inflammatory expression in breast cancer cells and induces anti-angiogenic, and anti-cancerous activities. Kahweol inhibits adipogenesis and increase glucose apoptosis in glioblastoma cells and lung cancer cells by activation of MEK-MAPK. Kaempferol can be uptake by AMP-activated protein kinase (AMPK) uesd for the research of breast cancer. activation. Kahweol induces apoptosis. Purity: 99 67% Purity: >98.0% Clinical Data: No Development Reported Clinical Data: No Development Reported 10 mM × 1 mL, 50 mg, 100 mg, 200 mg, 500 mg Size: Size: 5 mg, 10 mg, 20 mg Karanjin Kauran-16,17-diol Cat. No.: HY-N2534 (ent-Kauran-16β,17-diol) Cat. No.: HY-N7422 Karanjin is a major active furanoflavonol Kauran-16,17-diol (ent-Kauran-16β,17-diol), a constituent of Fordia cauliflora. Karanjin induces natural diterpene, posseses anti-tumor and GLUT4 translocation in skeletal muscle cells by inducing-apoptosis activity, with a IC_{50} of 17 μ M increasing AMPK activity. Karanjin can induce on inhibiting NO production in LPS-stimulated RAW cancer cell death through cell cycle arrest and 264.7 macrophages. enhance apoptosis. Purity: > 98% **Purity:** >98% Clinical Data: No Development Reported Clinical Data: No Development Reported 5 mg, 10 mg Size: Size: 1 ma **KEA1-97** Keap1-Nrf2-IN-4 Cat. No.: HY-114982 Cat. No.: HY-144099 KEA1-97 is a selective Thioredoxin-caspase 3 Keap1-Nrf2-IN-4 is a potent neddylation inhibitor. interaction disruptor (IC₅₀=10 μM). KEA1-97 Keap1-Nrf2-IN-4 exhibits potent anti-proliferation activity against MGC-803 cells (IC_{so} =2.55 µM). disrupts the interaction of thioredoxin with caspase 3, activates caspases, and induces Keap1-Nrf2-IN-4 blocks the migration ability and JAL D apoptosis without affecting thioredoxin activity. induces apoptosis of gastric cancer cells. 99.66% >98% Purity: Purity: Clinical Data: No Development Reported Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg Size: 1 mg, 5 mg Ketoprofen Ketoprofen-13C,d3 (RP-19583) (RP-19583-13C,d3) Cat. No.: HY-B0227 Cat. No.: HY-B0227S2 Ketoprofen (RP-19583) is a non-steroidal Ketoprofen-13C,d3 is the 13C- and deuterium labeled. Ketoprofen (RP-19583) is a non-steroidal antiinflammatory agent, acting as a potent inhibitor of COX, with IC₅₀s of 2 nM and 26 nM antiinflammatory agent, acting as a potent for COX-1 and COX-2 in human blood monocytes, inhibitor of COX, with IC50s of 2 nM and 26 nM for COX-1 and COX-2 in human blood monocytes, respectively. respectively. Purity: Purity: 99.93% >98% Clinical Data: Launched Clinical Data: No Development Reported 10 mM \times 1 mL, 500 mg, 1 g, 5 g Size: Size 1 mg, 5 mg Ketoprofen-d3 Ketoprofen-d4 (RP-19583-d3) Cat. No.: HY-B0227S (RP-19583-d4) Cat. No.: HY-B0227S1 Ketoprofen-d3 (RP-19583-d3) is the deuterium Ketoprofen-d4 (RP-19583-d4) is the deuterium labeled Ketoprofen. Ketoprofen (RP-19583) is a labeled Ketoprofen. Ketoprofen (RP-19583) is a non-steroidal antiinflammatory agent, acting as a non-steroidal antiinflammatory agent, acting as a potent inhibitor of COX, with IC₅₀s of 2 nM and potent inhibitor of COX, with IC50 s of 2 nM and 26 nM for COX-1 and COX-2 in human blood 26 nM for COX-1 and COX-2 in human blood monocytes, respectively. monocytes, respectively.

Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg



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Purity:

Size:

>98%

Clinical Data: No Development Reported

1 mg, 5 mg

Kinetin riboside		Kinsenoside	
(N6-Furfuryladenosine)	Cat. No.: HY-101055		Cat. No.: HY-N2292
Kinetin riboside, a cytokinin analog, can induce apoptosis in cancer cells. It inhibits the proliferation of HCT-15 cells with an IC_{s0} of 2.5 μ M.		Kinsenoside is a main active component isolated from plants of the genus Anoectochilus, and exhibits many biological activities and pharmacological effects.	
Purity:99.89%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 100 mg	HO	Purity:99.91%Clinical Data:No Development ReportedSize:5 mg, 10 mg	
KIRA9	Cat. No.: HY-145422	Kobe0065	Cat. No.: HY-15716
KIRA9 is a potent IRE1 inhibitor (IC_{50} =4.8 µM in INS-1 cells). KIRA9 is able to fully engage the ATP-binding site of IRE1 α . KIRA9 can block ER-localized mRNA decay and apoptosis .		Kobe0065 is a novel and effective inhibitor of Ras-Raf interaction , competitively inhibiting the binding of H-Ras-GTP to c-Raf-1 RBD with a K _i value of $46\pm13 \mu$ M.	
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	HÀ	Purity: 99.94% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg	9
Kongensin A		KP1019	
	Cat. No.: HY-N3417	(FFC14A)	Cat. No.: HY-19118
Kongensin A is a natural product isolated from Croton kongensis. Kongensin A is an effective, covalent HSP90 inhibitor that blocks RIP3-dependent necroptosishas. Kongensin A is a potent necroptosis inhibitor and an apoptosis inducer. Purity: ≥98.0% Clinical Data: No Development Reported Size: 1 mg, 5 mg	H H O H H H H	KP1019 (FFC14A) is a Ru(III)-based anti-metastatic and cytotoxic anti-cancer agent. KP1019 induces DNA damage and apoptosis in cancer cells. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	
KRAS G12D inhibitor 14	Cat. No.: HY-144661	KRAS inhibitor-9	Cat. No. : HY-137497
KRAS G12D inhibitor 14 is a potent KRAS G12D inhibitor with a K_p of 33 nM for binding to KRAS G12D protein. KRAS G12D inhibitor 14 decreases the active form of KRAS G12D (KRAS G12D-GTP) but not KRAS G13D.		KRAS inhibitor-9, a potent KRAS inhibitor (K_a =92 µM), blocks the formation of GTP-KRAS and downstream activation of KRAS. KRAS inhibitor-9 binds to KRAS G12D, KRAS G12C and KRAS Q61H protein with a moderate binding affinity.	
Clinical Data: No Development Reported Size: 1 mg, 5 mg		Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	00 mg
KRIBB11	Cat. No.: HY-100872	KS100	Cat. No.: HY-146682
KRIBB11 is an inhibitor of Heat shock factor 1 (HSF1), with IC _{s0} of 1.2 $\mu M.$		KS100 is a potent ALDH inhibitor with IC _{so} s of 230, 1542, 193 nM for ALDH1A1, ALDH2, and ALDH3A1, respectively. KS100 shows antiproliferative and anticancer effects with low low toxic.	
Purity: 99.12% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	NO ₂	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
KS106	Cat. No.: HY-146683	КТ5823	Cat. No.: HY-N6791
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KS106 is a potent ALDH inhibitor with IC ₅₀ s of 334, 2137, 360 nM for ALDH1A1, ALDH2, and ALDH3A1, respectively. KS106 shows antiproliferative and anticancer effects with low low toxic.	FLCH O HB	KT5823, a selective the cGMP-dependent protein kinase (PKG) inhibitor with an K ₁ value of 0.23 μ M, it also inhibits PKA and PKC with K ₁ values of 10 μ M and 4 μ M, respectively.	
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Purity:99.68%Clinical Data:No Development ReportedSize:100 μg	o N
Kumatakenin	Cat. No.: HY-N3415	Kurarinol	Cat. No.: HY-122933
Kumatakenin, a flavonoid that is isolated from cloves shows the effect of inducing apoptosis in ovarian cancer cells.	OF OF OF	Kurarinol is a flavanone found in the root of Sophora flavescens. Kurarinol is a competitive tyrosinase inhibitor, with IC ₅₀ of 0.1 μ M for mushroom tyrosinase.	HOT HO OH
Purity: ≥98.0% Clinical Data: No Development Reported Size: 5 mg, 10 mg		Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg	
KW-2449	Cat. No. : HY-10339	KY-05009	Cat. No .: HY-124745
KW-2449 is a multi-targeted kinase inhibitor of FLT3, ABL, ABL ^{T3151} and Aurora kinase with IC ₅₀ s of 6.6, 14, 4 and 48 nM, respectively.		KY-05009 is an ATP-competitive Traf2- and Nck-interacting kinase (TNIK) inhibitor with a K _i of 100 nM. KY-05009 pharmacologically inhibits TGF- β 1-induced epithelia1-to-mesenchymal transition (EMT) in human lung adenocarcinoma cells.	H2N NH HN S NH
Purity: 99.85% Clinical Data: Phase 1 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	OF M NH	Purity:99.80%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 25 mg, 100 mg	
Kynurenic acid (Quinurenic acid)	Cat. No.: HY-100806	Kynurenic acid sodium	Cat. No. : HY-107512
Kynurenic acid, an endogenous tryptophan metabolite, is a broad-spectrum antagonist targeting NMDA, glutamate, α? nicotinic acetylcholine receptor. Kynurenic acid is also an agonist of GPR35/CXCR8.	С ОН	Kynurenic acid sodium, an endogenous tryptophan metabolite, is a broad-spectrum antagonist targeting NMDA, glutamate, α7 nicotinic acetylcholine receptor. Kynurenic acid sodium is also an agonist of GPR35/CXCR8.	ONa
Purity: 99.58% Clinical Data: Phase 1 Size: 10 mM × 1 mL, 100 mg, 500 mg	ОН	Purity: 99.76% Clinical Data: Phase 1 Size: 10 mM × 1 mL, 100 mg	ОН
Kynurenic acid-d5 (Quinurenic acid-d5)	Cat. No.: HY-100806S	КҮР-2047	Cat. No.: HY-100475
Kynurenic acid-d5 (Quinurenic acid-d5) is the deuterium labeled Kynurenic acid. Kynurenic acid, an endogenous tryptophan metabolite, is a broad-spectrum antagonist targeting NMDA, glutamate, α7 nicotinic acetylcholine receptor.</b 		KYP-2047 is a potent and BBB-penetrating prolyl-oligopeptidase (POP) inhibitor, with an K _i value of 0.023 nM. KYP-2047 reduces glioblastoma proliferation through angiogenesis and apoptosis modulation.	
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg, 25 mg		Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	V 0

L-685458		L-Ascorbic acid	
(L-685,458)	Cat. No.: HY-19369	(L-Ascorbate; Vitamin C)	Cat. No.: HY-B0166
L-685458 is a potent transition state analog (TSA) γ -secretase inhibitor (GSI). L-685458 inhibits amyloid β -protein precursor γ -secretase activity with IC _{so} of 17 nM, shows greater than 50-100-fold selectivity over other aspartyl proteases tested.	July Contractions	L-Ascorbic acid (L-Ascorbate), an electron donor, is an endogenous antioxidant agent. L-Ascorbic acid inhibits selectively Ca_v3.2 channels with an IC _{so} of 6.5 μ M. L-Ascorbic acid is also a collagen deposition enhancer and an elastogenesis inhibitor.	HO HO OH
Purity: 99.33%		Purity: 99.92%	
Clinical Data: No Development Reported		Clinical Data: Launched	
Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg		Size: 10 mM × 1 mL, 500 mg, 1 g	
L-Ascorbic acid sodium salt		L-Ascorbic acid-13C	
(Sodium L-ascorbate; Vitamin C sodium salt)	Cat. No.: HY-B0166A	(L-Ascorbate-13C; Vitamin C-13C)	Cat. No.: HY-B0166S1
L-Ascorbic acid sodium salt (Sodium L-ascorbate), an electron donor, is an endogenous antioxidant agent. L-Ascorbic acid sodium salt inhibits selectively Ca_v3.2 channels with an IC ₅₀ of 6.5 μ M.		L-Ascorbic acid-13C (L-Ascorbate-13C) is the 13C-labeled L-Ascorbic acid. L-Ascorbic acid (L-Ascorbate), an electron donor, is an endogenous antioxidant agent. L-Ascorbic acid inhibits selectively Ca _v 3.2 channels with an IC ₅₀ of 6.5 µM.	но он 012 он
Purity: 99.17% Clinical Data: Launched		Purity: >98% Clinical Data: No Development Reported	
Size: 10 mM × 1 mL, 500 mg, 1 g		Size: 1 mg, 5 mg	
L-Ascorbic acid-13C6		L-Cystathionine	
(L-Ascorbate-13C6; Vitamin C-13C6)	Cat. No.: HY-B0166S		Cat. No.: HY-W009749
L-Ascorbic acid-13C6 (L-Ascorbate-13C6) is the 13C-labeled L-Ascorbic acid. L-Ascorbic acid (L-Ascorbate), an electron donor, is an endogenous antioxidant agent. L-Ascorbic acid inhibits selectively Ca _v 3.2 channels with an IC _{so} of 6.5 μ M. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	93С-0 Н2 13С-13С-11-13С 132 H0 13C-14-13С H0 0H ОН 0H	L-Cystathionine is a nonprotein thioether and is a key amino acid associated with the metabolic state of sulfur-containing amino acids. L-Cystathionine protects against Homocysteine-induced mitochondria-dependent apoptosis of vascular endothelial cells (HUVECs). Purity: ≥98.0% Clinical Data: No Development Reported Size: 5 mg, 10 mg	
L Clutamic acid		L. Glutamic acid monocodium calt	
	Cat No : HY-14608		Cat No: HY-14608A
L-Glutamic acid acts as an excitatory transmitter and an agonist at all subtypes of glutamate receptors (metabotropic, kainate, NMDA, and AMPA). L-Glutamic acid shows a direct activating effect on the release of DA from dopaminergic terminals. Purity: ≥98.0% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 500 mg	H ₂ N OH	L-Glutamic acid monosodium salt acts as an excitatory transmitter and an agonist at all subtypes of glutamate receptors (metabotropic, kainate, NMDA, and AMPA). (S)-Glutamic acid shows a direct activating effect on the release of DA from dopaminergic terminals. Purity: $\geq 98.0\%$ Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 500 mg	H ₂ N OH
L-Glutamic acid-1-13C	Cat. No.: HY-14608S1	L-Glutamic acid-13C	Cat. No.: HY-14608S
L-Glutamic acid-1-13C is the 13C-labeled L-Glutamic acid. L-Glutamic acid acts as an excitatory transmitter and an agonist at all subtypes of glutamate receptors (metabotropic, kainate, NMDA, and AMPA).	но о NH ₂	L-Glutamic acid-13C is the 13C-labeled L-Glutamic acid. L-Glutamic acid acts as an excitatory transmitter and an agonist at all subtypes of glutamate receptors (metabotropic, kainate, NMDA, and AMPA).	HO H2 OH NH2 OH
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	

L-Glutamic acid-13C5	Cat. No.: HY-14608S5	L-Glutamic acid-13C5,15N	Cat. No. : HY-14608S3
L-Glutamic acid-13C5 is the 13C-labeled L-Glutamic acid. L-Glutamic acid acts as an excitatory transmitter and an agonist at all subtypes of glutamate receptors (metabotropic, kainate, NMDA, and AMPA).	О H2 О H0 ¹³ G3C ⁻¹³ G ¹³ C H0 ¹³ G3C ⁻⁰ H H2 NH2	L-Glutamic acid-13C5,15N is the 13C- and 15N-labeled L-Glutamic acid. L-Glutamic acid acts as an excitatory transmitter and an agonist at all subtypes of glutamate receptors (metabotropic, kainate, NMDA, and AMPA).	о но но но на на на на на на на на на на на на на
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	
L-Glutamic acid-13C5,d5,15N	Cat. No.: HY-14608S4	L-Glutamic acid-15N	Cat. No. : HY-14608S2
L-Glutamic acid-13C5,d5,15N is the deuterium, 13C-, and 15-labeled L-Glutamic acid. L-Glutamic acid acts as an excitatory transmitter and an agonist at all subtypes of glutamate receptors (metabotropic, kainate, NMDA, and AMPA).	О D D O но ¹³ СзС ¹³ Сз _С ¹³ С-он пн₂ы D	L-Glutamic acid-15N is the 15N-labeled L-Glutamic acid. L-Glutamic acid acts as an excitatory transmitter and an agonist at all subtypes of glutamate receptors (metabotropic, kainate, NMDA, and AMPA).	но о о 15 _{NH2} он
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:>98%Clinical Data:No Development ReportedSize:25 mg, 50 mg, 100 mg	
L-Glutamic acid-15N,d5	Cat. No.: HY-14608S9	L-Glutamic acid-5-13C	Cat. No. : HY-14608S6
L-Glutamic acid-15N,d5 is the deuterium and 15N-labeled L-Glutamic acid. L-Glutamic acid acts as an excitatory transmitter and an agonist at all subtypes of glutamate receptors (metabotropic, kainate, NMDA, and AMPA). Purity: >98% Clinical Data: No Development Reported	$H_2^{15}N_4$ OH OH OH	L-Glutamic acid-5-13C is the 13C-labeled L-Glutamic acid. L-Glutamic acid acts as an excitatory transmitter and an agonist at all subtypes of glutamate receptors (metabotropic, kainate, NMDA, and AMPA). Purity: >98% Clinical Data: No Development Reported	HO ¹³ C OH NH ₂ OH
Size: 1 mg, 5 mg		Size: 1 mg, 5 mg	
L-Glutamic acid-d3	Cat. No. : HY-14608S8	L-Glutamic acid-d5	Cat. No. : HY-14608S7
L-Glutamic acid-d3 is the deuterium labeled L-Glutamic acid. L-Glutamic acid acts as an excitatory transmitter and an agonist at all subtypes of glutamate receptors (metabotropic, kainate, NMDA, and AMPA).		L-Glutamic acid-d5 is the deuterium labeled L-Glutamic acid. L-Glutamic acid acts as an excitatory transmitter and an agonist at all subtypes of glutamate receptors (metabotropic, kainate, NMDA, and AMPA).	
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
L-SelenoMethionine	Cat. No.: HY-B1000A	L-Theanine (L-Glutamic Acid y-ethyl amide; Ny-Ethyl-L-glutamine)	Cat. No.: HY-15121
L-SelenoMethionine, an L-isomer of Selenomethionine, is a major natural food-form of selenium. L-SelenoMethionin is a cancer chemopreventive agent that can reduce cancer incidence by dietary supplementation and induce apoptosis of cancer cells. Purity: 99.84% Clinical Data: Launched	Se OH NH2	L-Theanine (L-Glutamic Acid γ-ethyl amideis a non-protein amino acid contained in green tea leaves, which blocks the binding of L-glutamic acid to glutamate receptors in the brain, and with neuroprotective and anti-oxidative activities. Purity: 98.84% Clinical Data: Phase 4	
Size: 10 mM × 1 mL, 100 mg, 500 mg		Size: 10 mM × 1 mL, 100 mg, 200 mg	

L-Theanine-d5 (L-Glutamic Acid γ-ethyl amide-d5;		L-threo-PPMP	
Nγ-Ethyl-L-glutamine-d5)	Cat. No.: HY-15121S		Cat. No.: HY-115737
L-Theanine-d5 (L-Glutamic Acid γ-ethyl amide-d5) is the deuterium labeled L-Theanine. Purity: >98%		L-threo-PPMP is a GlcT (UDP-Glc: Ceramide β1,1glucosyltransferase) inhibitor. L-threo-PPMP inhibits glycosphingolipid biosynthesis and induces apoptosis. L-threo-PPMP has anti-cancer activity.	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
Clinical Data: No Development Reported Size: 1 mg, 5 mg		Clinical Data: No Development Reported Size: 1 mg, 5 mg	
L5-DA	Cat. No.: HY-144712	Lacidipine	Cat. No.: HY-B0347
L5-DA is a G-quadruplex (G4) ligand and selectively stabilized for G4s over ds26. L5-DA exhibits significant cytotoxicity against HeLa cells (IC ₅₀ =4.3 μ M). L5-DA stabilizes G4s in HeLa cells, induces apoptosis , and cell cycle arrest.	4.09000.+	Lacidipine (Lacipil, Motens) is a L-type calcium channel blocker. Target: Calcium Channel Lacidipine, a novel third-generation dihydropyridine calcium channel blocker, has been demonstrated effective for hypertension.	
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Purity: 99.98% Clinical Data: Launched Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg	
Lacidipine-d10	Cat. No.: HY-B0347S	Lactoferrin (17-41) (Lactoferricin B; Lfcin B)	Cat. No. : HY-P1791
Lacidipine-d10 is the deuterium labeled Lacidipine. Lacidipine (Lacipil, Motens) is a L-type calcium channel blocker.		Lactoferrin 17-41 (Lactoferricin B), a peptide corresponding to residues 17-41 of bovine lactoferrin, has antimicrobial activity against a wide range of microorganisms, including Gram-positive and Gramnegative bacteria, viruses, protozoa, and fungi.	псяворнаясынатсяная рыла шар сулсурст
Clinical Data: Size: 1 mg, 10 mg		Clinical Data: No Development Reported Size: 1 mg, 5 mg	
Lactoferrin (17-41) (acetate) (Lactoferricin B acetate; Lfcin B acetate)	Cat. No.: HY-P1791B	Lactonic sophorolipid	Cat. No.: HY-137371
Lactoferrin 17-41 (Lactoferricin B) acetate, a peptide corresponding to residues 17-41 of bovine lactoferrin, has antimicrobial activity against a wide range of microorganisms, including Gram-positive and Gramnegative bacteria, viruses, protozoa, and fungi.	FICORRAQUEIMOL QAPSTCAREAF (Disufter Entge: Cyc3-Cyc20) (soviste sal)	Lactonic sophorolipid is a natural antimicrobial surfactant for oral hygiene. Lactonic sophorolipid, a potential anticancer agent, induces apoptosis in human HepG2 cells through the caspase-3 pathway.	
Clinical Data: No Development Reported Size: 5 mg, 10 mg		Clinical Data: No Development Reported Size: 1 mg, 5 mg	
Laquinimod (ABR-215062)	Cat. No.: HY-13010	Laquinimod sodium (ABR-215062 sodium)	Cat. No.: HY-W062904
Laquinimod (ABR-215062), an orally available carboxamide derivative, is a potent immunomodulator which prevents neurodegeneration and inflammation in the central nervous system.		Laquinimod (ABR-215062) sodium, an orally available carboxamide derivative, is a potent immunomodulator which prevents neurodegeneration and inflammation in the central nervous system.	CI OH O
Purity: 99.91% Clinical Data: Launched Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg,	500 mg	Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	



Leachianone A		Lenalidomide	
	Cat. No.: HY-N2281	(CC-5013)	Cat. No.: HY-A0003
Leachianone A, isolated from Radix Sophorae, has anti-malarial, anti-inflammatory, and cytotoxic potent. Leachianone A induces apoptosis involved both extrinsic and intrinsic pathways	HO CO O' OH	Lenalidomide (CC-5013), a derivative of Thalidomide, acts as molecular glue. Lenalidomide is an orally active immunomodulator.	
Purity: ≥98.0% Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg		Purity: 99.91% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 500 mg, 1 g	
Lenalidomide hemihydrate (CC-5013 hemihydrate)	Cat. No.: HY-A0003B	Lenalidomide-d5 (CC-5013-d5)	Cat. No.: HY-A0003S
Lenalidomide hemihydrate (CC-5013 hemihydrate), a derivative of Thalidomide, acts as molecular glue. Lenalidomide hemihydrate is an orally active immunomodulator.		Lenalidomide-d5 is deuterium labeled Lenalidomide. Lenalidomide (CC-5013), a derivative of Thalidomide, acts as molecular glue. Lenalidomide is an orally active immunomodulator.	
Purity: 99.95% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 500 mg	0.5 H ₂ O	Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	
Lepidozin G	Cat. No. : HY-141863	Levistolide A	Cat. No.: HY-N1472
Lepidozin G inhibits the growth of a panel of cancer cell lines with IC_{s0} values ranging from 4.2 ± 0.2 to 5.7 ± 0.5 μ M. Lepidozin G induces PC-3 cell death via mitochondrial-related apoptosis .	HO TH IN TO OH	Levistolide A (LA), a natural compound isolated from the traditional Chinese herb Ligusticum chuanxiong Hort, is used for treating cancer. Levistolide A can induce apoptosis via ROS-mediated ER stress pathway.	
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:98.87%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 20 mg	O H
Levomenol ((-)-α-Bisabolol)	Cat No : HY-N6967	Lexibulin (CYT-997)	Cat No : HV-10498
Levomenol ((-)- α -Bisabolol), a monocyclic sesquiterpene alcohol, exerts antioxidant, anti-inflammatory, and anti-apoptotic activities.		Lexibulin (CYT-997) is a potent and orally active tubulin polymerisation inhibitor with IC50s of 10-100 nM in cancer cell lines; with potent cytotoxic and vascular disrupting activity in vitro and in vivo.	
Purity:98.35%Clinical Data:No Development ReportedSize:5 mL		Purity: 98.08% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg	C [™]
Lexibulin dihydrochloride (CYT-997 dihydrochloride)	Cat. No. : HY-10498A	LG308	Cat. No.: HY-143660
Lexibulin dihydrochloride (CYT-997 dihydrochloride) is a potent and orally active tubulin polymerisation inhibitor with IC50s of 10-100 nM in cancer cell lines; with potent cytotoxic and vascular disrupting activity in vitro and in vivo.		LG308 is a novel synthetic compound with antimicrotubule activity. LG308 induces mitotic phase arrest and inhibits G2/M progression significantly which is associated with the upregulation of cyclin B1 and mitotic marker MPM-2 and the dephosphorylation of cdc2.	5000
Purity:>98%Clinical Data:Phase 2Size:1 mg, 5 mg	ι, Ν.	Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	

Licochalcone B	Cat. No.: HY-N0373	Licofelone (ML-3000)	Cat. No.: HY-B1452
Licochalcone B is an extract from the root of Glycyrrhiza inflate.	но он	Licofelone (ML-3000) is a dual COX/5-lipoxygenase (5-LOX) inhibitor (IC ₅₀ =0.21/0.18 μ M, respectively) for the treatment of osteoarthritis. Licofelone exerts anti-inflammatory and anti-proliferative effects.	с-
Purity:99.93%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 1 mg, 5 mg, 10 mg		Purity:98.04%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 1 mg, 5 mg, 10 mg	
Licofelone-d4	Cat. No.: HY-B1452S	Licoricidin	Cat. No.: HY-N3387
Licofelone-d4 (ML-3000-d4) is the deuterium labeled Licofelone. Licofelone (ML-3000) is a dual COX/5-lipoxygenase (5-LOX) inhibitor (IC_{50} =0.21/0.18 µM, respectively) for the treatment of osteoarthritis.		Licoricidin (LCD) is isolated from Glycyrrhiza uralensis Fisch, possesses anti-cancer activities.	HO, COH
Purity:>98%Clinical Data:Size:5 mg		Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg	
Lideraine		Lide seine kudus sklevide	
(Lignocaine)	Cat. No.: HY-B0185	(Lignocaine hydrochloride)	Cat. No.: HY-B0185A
Lidocaine (Lignocaine) inhibits sodium channels involving complex voltage and using dependence.		Lidocaine hydrochloride (Lignocaine hydrochloride) inhibits sodium channels involving complex voltage and using dependence.	L H N
Purity: 99.96% Clinical Data: Launched Size: 10 mM × 1 mL, 500 mg, 5 g, 10 g		Purity: 99.81% Clinical Data: Launched Size: 10 mM × 1 mL, 500 mg, 5 g, 10 g	H-CI
Lidocaine-d10	Cat. No.: HY-B0185S1	Lidocaine-d10 hydrochloride	Cat. No.: HY-B0185AS
Lidocaine-d10 is the deuterium labeled Lidocaine. Lidocaine (Lignocaine) inhibits sodium channels involving complex voltage and using dependence.		Lidocaine-d10 (Lignocaine-d10) hydrochloride is the deuterium labeled Lidocaine hydrochloride. Lidocaine hydrochloride (Lignocaine hydrochloride) inhibits sodium channels involving complex voltage and using dependence.	
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 50 mg	р р нсі
Lidocaine-d10 N-Oxide	Cat. No.: HY-B0185S	Lidocaine-d6 hydrochloride (Lignocaine-d6 hydrochloride)	Cat. No.: HY-B0185AS1
Lidocaine-d10 N-Oxide is the deuterium labeled Lidocaine. Lidocaine (Lignocaine) inhibits sodium channels involving complex voltage and using dependence.		Lidocaine-d6 (hydrochloride) is deuterium labeled Lidocaine (hydrochloride). Lidocaine hydrochloride (Lignocaine hydrochloride) inhibits sodium channels involving complex voltage and using dependence.	
Purity:>98%Clinical Data:No Development ReportedSize:2.5 mg, 25 mg	D	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	D



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Lithocholic acid-d4		Lithocholic acid-d5	
(3α-Hydroxy-5β-cholanic acid-d4)	Cat. No.: HY-B0172S	(3α-Hydroxy-5β-cholanic acid-d5)	Cat. No.: HY-B0172S1
Lithocholic acid-d4 (3α-Hydroxy-5β-cholanic acid-d4) is the deuterium labeled Lithocholic acid, which is a toxic secondary bile acid.		Lithocholic acid-d5 is deuterium labeled Lithocholic acid.	
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg, 10 mg, 25 mg		Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	
LJH685	Cat. No. : HY-19712	Lobetyolin	Cat. No.: HY-N0327
LJH685 is a potent, ATP-competitive and selective RSK inhibitor, inhibits RSK1, 2, and 3 biochemical activities with IC ₅₀ s of 6, 5, 4 nM, respectively. Purity: 99.99% Clinical Data: No Development Reported		Lobetyolin, a bioactive compound, is derived from Codonopsis pilosula. Lobetyolin has anti-inflammatory, anti-oxidative and xanthine oxidase inhibiting activities. Lobetyolin also induces the apoptosis via the inhibition of ASCT2-mediated glutamine metabolism. Purity: 99.89% Clinical Data: No Development Reported	HO CH HO CH HO CH
Size: $10 \text{ mM} \times 1 \text{ mL}$, 5 mg, 10 mg, 50 mg, 100 mg		Size: 5 mg	
5122. 20 mm 8 2 m2, 5 mg, 20 mg, 50 mg, 200 mg		5.120. 5 mg	
Loganin		Lometrexol	
(Loganoside)	Cat. No.: HY-N0512	(DDATHF)	Cat. No.: HY-14521
Loganin, a major iridoid glycoside obtained from Corni fructus, has been shown to have anti-inflammatory and anti-shock effects. Loganin exhibits an anti-inflammatory effect in cases of AP and its pulmonary complications through inhibition of NF-κB activation.		Lometrexol (DDATHF), an antipurine antifolate , can inhibit the activity of glycinamide ribonucleotide formyltransferase (GARFT) but do not induce detectable levels of DNA strand breaks.	
Purity: 99.82% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg	O O	Purity: >98% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg	
Lometrexol hydrate		Lomustine	
(DDATHF hydrate)	Cat. No.: HY-14521B	(CCNU; NSC 79037)	Cat. No.: HY-13669
Lometrexol hydrate (DDATHF hydrate), an antipurine antifolate, can inhibit the activity of glycinamide ribonucleotide formyltransferase (GARFT) but do not induce detectable levels of DNA strand breaks.	What has no	Lomustine (CCNU; NSC 79037) is a DNA alkylating agent, with antitumor activity.	H N ² O N ² O N ² O
Purity:99.20%Clinical Data:No Development ReportedSize:5 mg, 10 mg		Purity: 99.91% Clinical Data: Launched Size: 10 mM × 1 mL, 200 mg, 500 mg	
Lonicerin	Cat. No.: HY-N4136	Lonidamine (AF-1890; Diclondazolic Acid; DICA)	Cat. No.: HY-B0486
Lonicerin is an anti- algE (alginate secretion protein) flavonoid with inhibitory activity for P. aeruginosa. Lonicerin prevents inflammation and apoptosis in LPS-induced acute lung injury.		Lonidamine (AF-1890), an antitumor agent, is a hexokinase, mitochondrial pyruvate carrier (K_i 2.5 μ M in isolated rat liver mitochondria) and plasma membrane monocarboxylate transporters inhibitor, which also inhibits mitochondrial complex II.	
Purity:99.75%Clinical Data:No Development ReportedSize:5 mg, 10 mg		Purity: 99.45% Clinical Data: Phase 3 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg	0′ 0′



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Luteolin		LW6	
Luteoline; Luteoloi; Digitofiavone) Luteolin (Luteoline), a flavanoid compound, is a potent Nrf2 inhibitor.		(HIF-1α inhibitor) is a novel HIF-1 inhibitor with an IC ₅₀ of 4.4 μM. LW6 decreases HIF-1α protein expression without affecting HIF-1β expression.	
Purity: 98.42% Clinical Data: Phase 1 Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 500 mg		Purity: 98.93% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	10 mg
LY-411575	Cat. No. : HY-50752	LY2409881 trihydrochloride	Cat. No.: HY-B0788A
LY-411575 is a potent γ -secretase inhibitor with IC ₅₀ of 0.078 nM/0.082 nM (membrane/cell-based), and also inhibits Notch S3 cleavage with IC ₅₀ of 0.39 nM.		LY2409881 trihydrochloride is a selective I κ B kinase β (IKK2) inhibitor with an IC ₅₀ of 30 nM.	HW O H-CO V
Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	in the second s	Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
LY2857785	Cat. No.: HY-12293	LY294002	Cat. No. : HY-10108
LY2857785 is a type I reversible and competitive ATP kinase inhibitor against CDK9 (IC_{s0} 11 nM) and other transcription kinases CDK8 (IC_{s0} 16 nM), and CDK7 (IC_{s0} 246 nM).		LY294002 is a broad-spectrum inhibitor of PI3K with IC_{so} of 0.5, 0.57, and 0.97 μ M for $PI3K\alpha$, $PI3K\delta$ and $PI3K\beta$, respectively. LY294002 also inhibits CK2 with an IC_{so} of 98 nM.	
Purity:98.88%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 50 mg, 100 mg	g	Purity: 99.95% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg	o 9, 500 mg
Lycopodine	C at. No.: HY-114372	Lycorine	Cat. No.: HY-N0288
Lycopodine, a pharmacologically important bioactive component derived from Lycopodium clavatumspores, triggers apoptosis by modulating 5-lipoxygenase , and depolarizing mitochondrial membrane potential in refractory prostate cancer cells without modulating p53 activity.		Lycorine is a natural alkaloid extracted from the Amaryllidaceae plant. Lycorine is a potent and orally active SCAP inhibitor with a K_a value 15.24 nM. Lycorine downregulates the SCAP protein level without changing its transcription.	
Clinical Data: No Development Reported Size: 1 mg, 5 mg	\checkmark	Clinical Data: No Development Reported Size: 50 mg, 100 mg	
LYN-1604	C at. No.: HY-101923	LYN-1604 dihydrochloride	Cat. No.: HY-101923B
LYN-1604 is a potent UNC-51-like kinase 1 (ULK1) activator (EC_{s0} =18.94 nM) for the research of triple negative breast cancer (TNBC).		LYN-1604 dihydrochloride is a potent UNC-51-like kinase 1 (ULK1) activator (EC_{so} =18.94 nM) for the research of triple negative breast cancer (TNBC).	
Purity: >98% c Clinical Data: No Development Reported Size: 1 mg, 5 mg		Purity: 98.73% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	H-Ci H-Ci



Maduramicin ammonium (Maduramycin ammonium)	Cat. No. : HY-N7071A	Malachite green oxalate	Cat. No.: HY-D0162
Maduramicin ammonium (Maduramycin ammonium) is isolated from the actinomycete Actinomadura rubra. Purity: ≥98.0%		Malachite green oxalate is a triphenylmethane dye which can be used to detect the release of phosphate in enzymatic reactions. Malachite green oxalate is also a potent and selective inhibitor of IKBKE, and inhibits its downstream targets such as IkB α , p65 and IRF3. Purity: \geq 98.0%	^{- Н} ОДО ^{- К} юдо 12 юдон
Clinical Data: No Development Reported Size: 5 mg, 10 mg		Clinical Data: No Development Reported Size: 10 mM × 1 mL, 100 mg	
Malformin A1	Cat. No.: HY-P2569	Malvidin-3-galactoside chloride	Cat. No.: HY-N6623
Malformin A1, a cyclic pentapeptide isolated from Aspergillus niger, possess a range of bioactive properties including antibacterial activity. Malformin A1 shows potent cytotoxic activities on human colorectal cancer cells.		Malvidin-3-galactoside chloride, an anthocyanin monomer, induces hepatocellular carcinoma (HCC) cells cycle arrest and apoptosis . Malvidin-3-galactoside chloride inhibits the production and accumulation of ROS .	
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	0	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg, 10 mg	он
Mangiferin		Mangostin-d3	
	Cat. No.: HY-N0290		Cat. No.: HY-N0328S
Mangiferin is a Nrf2 activator. Mangiferin suppresses nuclear translocation of the NF-κB subunits p65 and p50 . Mangiferin exhibits antioxidant, antidiabetic, antihyperuricemic, antiviral, anticancer and antiinflammatory activities.		alpha-Mangostin-d3 (α -Mangostin-d3) is the deuterium labeled alpha-Mangostin. alpha-Mangostin (α -Mangostin) is a dietary xanthone with broad biological activities, such as antioxidant, anti-allergic, antiviral, antibacterial, anti-inflammatory and anticancer effects.	Hold of the
Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg		Clinical Data: Size: 2.5 mg, 25 mg	
Manumvcin A		MARK4 inhibitor 1	
	Cat. No.: HY-N6796		Cat. No.: HY-114317
Manumycin A is an antibiotic. Manumycin A acts as a selective, competitive inhibitor of protein farnesyltransferase (FTase) with respect to farnesylpyrophosphate ($K_i = 1.2 \mu$ M), and as a noncompetitive inhibitor with respect to the Ras protein.		MARK4 inhibitor 1 is a potent microtubule affinity-regulating kinase 4 (MARK4) inhibitor, with an IC_{so} of 1.54 μ M. MARK4 inhibitor 1 inhibits cancer cell proliferation, metastasis and induces apoptosis.	$ \begin{array}{c} \begin{array}{c} & & \\ & & \\ & & \\ & & \\ & & \\ & HN \\ & HN \\ & HN \\ & HN \\ & \\ & HN \\ & \\ & HN \\ & \\ & \\ & \\ & \\ & \\ & \\ & \\ & \\ & \\ $
Clinical Data: No Development Reported Size: 1 mg, 5 mg		Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	10 mg
Masitinib		Masitinib mesylate	
(AB1010)	Cat. No.: HY-10209	(AB-1010 mesylate)	Cat. No.: HY-10209A
Masitinib (AB1010) is a potent, orally bioavailable, and selective inhibitor of c-Kit (IC_{so} =200 nM for human recombinant c-Kit). It also inhibits PDGFRa (β (IC_{so} S=540/800 nM), Lyn (IC_{so} =510 nM for LynB), Lck, and, to a lesser extent, FGFR3 and FAK.	watara	Masitinib mesylate (AB-1010 mesylate) is a potent, orally bioavailable, and selective inhibitor of c-Kit (IC _{so} =200 nM for human recombinant c-Kit). It also inhibits PDGFR α/β (IC _{so} s=540/800 nM), Lyn (IC _{so} = 510 nM for LynB), Lck, and, to a lesser extent, FGFR3 and FAK.	in and the second
Purity: 99.98% Clinical Data: Phase 3 Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg	I	Purity: 99.76% Clinical Data: Phase 3 Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg]



MDK83190MebendazoleCat. No: HY-18633Cat. No: HY-18633MDK83190 is a potent apoptosis activator, induces Apaf-1 oligomerization, increases procaspase-9 processing and subsequent caspase-3 activation in a cyto c-dependent Manner. $\qquad \qquad $
MDK83190 is a potent apoptosis activator, induces Apaf-1 oligomerization, increases procaspase-9 processing and subsequent caspase-3 activation in a cyto c-dependent Manner.Mebendazole is a highly effective, broad-spectrum antiblelmintic indicated for the treatment of nematode infestations, has been found as a hedgehog inhibitor.Mebendazole is a highly effective, broad-spectrum antiblelmintic indicated for the treatment of nematode infestations, has been found as a hedgehog inhibitor.Mebendazole-d8Cat. No: HY-1759551Mechercharmycin A s a cytotoxic substance isolated for marine-derived Thermoactinomyces sp. YM3-251. Mechercharmycin A exhibits relatively strong antitumor activity.Cat. No: HY-13625Metodicarpin $\mu_{p}^{+} + \mu_{p}^{+} + \mu_{p$
Purity:98.08% Clinical Data:Purity:99.88% Clinical Data:Mebendazole-d8Cat. No:: HY-1759551Mechercharmycin AMebendazole-d8 is the deuterium labeled Mebendazole. Mebendazole is a highly effective, broad-spectrum antihelmintic indicated for the treatment of nematode infestations; has been found as a hedgehog inhibitor. $p + f + f + f + f + f + f + f + f + f + $
Mebendazole-d8 Cat. No.: HY-1759551 Mechercharmycin A Cat. No.: HY-13625 Mebendazole. d8 is the deuterium labeled Mebendazole. Mebendazole is a highly effective, broad-spectrum antihelmintic indicated for the treatment of nematode infestations; has been found as a hedgehog inhibitor. Image: Comparison of the treatment of nematode infestations; has been found as a hedgehog inhibitor. Image: Comparison of the treatment of nematode infestations; has been found as a hedgehog inhibitor. Image: Comparison of the treatment of nematode infestations; has been found as a hedgehog inhibitor. Image: Comparison of the treatment of nematode infestations; has been found as a hedgehog inhibitor. Image: Comparison of the treatment of nematode infestations; has been found as a hedgehog inhibitor. Image: Comparison of the treatment of nematode infestations; has been found as a hedgehog inhibitor. Image: Comparison of the treatment of nematode infestations; has been found as a hedgehog inhibitor. Image: Comparison of the treatment of nematode infestations; has been found as a hedgehog inhibitor. Image: Comparison of the treatment of nematode infestations; has been found as a hedgehog inhibitor. Image: Comparison of the treatment of nematode infestations; has been found as a hedgehog inhibitor. Image: Comparison of the treatment of nematode infestations; has been found as a hedgehog inhibitor. Image: Comparison of the treatment of nematode infestations; has been found as a hedgehog inhibitor. Image: Comparison of the treatment of nematode infestations; has been found as a hedgehog inhibitor. Image: Comparison of the treatment of nematode infestations; has been found as a hedgehog inhibitor. <
Mebendazole-d8 is the deuterium labeled Mebendazole. Mebendazole is a highly effective, broad-spectrum antihelmintic indicated for the treatment of nematode infestations; has been found as a hedgehog inhibitor. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg Medicarpin Mefuparib hydrochloride
Purity: 598% Clinical Data: No Development Reported Size: 1 mg, 5 mg Medicarpin Mefuparib hydrochloride
Medicarpin Mefuparib hydrochloride
Cat. No.: HY-N3308 (MPH) Cat. No.: HY-12266
Medicarpin is a flavonoid isolated from Medicago sativa. Medicarpin induces apoptosis and overcome multidrug resistance in leukemia P388 cells by modulating P-gp-mediated efflux of drugs.Mefuparib hydrochloride (MPH) is an orally active, substrate-competitive and selective PARP1/2 inhibitor with IC ₅₀ s of 3.2 nM and 1.9 nM, respectively. Mefuparib hydrochloride induces apoptosis and possesses prominent anticancer activity in vitro and in vivo. Purity: \geq 98.0% Clinical Data: No Development ReportedPurity: No Development Reported
Size: 1 mg, 5 mg Size: 5 mg, 10 mg, 25 mg
Meisoindigo MEK-IN-5 (Dian III; N-Methylisoindigotin; Natura-α) Cat. No.: HY-13680
Meisoindigo (Dian III), a derivative of Indirubin (HY-N0117), halts the cell cycle at the G0/G1 phase and induces apoptosis in primary acute myeloid leukemia (AML) cells. Meisoindigo exhibits high antitumor activity. Purity: 98.08% Clinic D is to be
Clinical Data: Launched Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg Size: 1 mg, 5 mg
Melatonin Melatonin-d3 (N-Acetyl-5-methoxytryptamine) Cat. No.: HY-B0075 (N-Acetyl-5-methoxytryptamine-d3) Cat. No.: HY-B00755
Melatonin is a hormone made by the pineal gland that can activates melatonin receptor . Melatonin plays a role in sleep and possesses important antioxidative and anti-inflammatory properties.
Purity: 99.73% Purity: >98% Clinical Data: Launched Clinical Data: No Development Reported Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g Size: 1 mg, 5 mg

Melatonin-d4		Melatonin-d7	
(N-Acetyl-5-methoxytryptamine-d4)	Cat. No.: HY-B0075S	(N-Acetyl-5-methoxytryptamine-d7)	Cat. No.: HY-B0075S2
Melatonin D4 is deuterium labeled Melatonin. Melatonin is a hormone made by the pineal gland that can activates melatonin receptor. Antioxidative and anti-inflammatory properties.		Melatonin-d7 (N-Acetyl-5-methoxytryptamine-d7) is the deuterium labeled Melatonin. Melatonin is a hormone made by the pineal gland that can activates melatonin receptor .	
Purity:95.87%Clinical Data:No Development ReportedSize:1 mg, 5 mg, 10 mg	↓	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg, 10 mg	
Melflufen		Melflufen hydrochloride	
(Melphalan flufenamide)	Cat. No.: HY-105019	(Melphalan flufenamide hydrochloride)	Cat. No.: HY-105019A
Melflufen (Melphalan flufenamide), a dipeptide prodrug of Melphalan, is an alkylating agent. Melflufen shows antitumor activity against multiple myeloma (MM) cells and inhibits angiogenesis. Melflufen induces irreversible DNA damage and cytotoxicity in MM cells.		Melflufen (Melphalan flufenamide) hydrochloride, a dipeptide prodrug of Melphalan, is an alkylating agent. Melflufen hydrochloride shows antitumor activity against multiple myeloma (MM) cells and inhibits angiogenesis.	
Purity: >98% Clinical Data: Launched		Purity: 99.20% Clinical Data: Launched	
Size: 1 mg, 5 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
Meloxicam		Meloxicam-13C,d3	
	Cat. No.: HY-B0261		Cat. No.: HY-B0261S2
Meloxicam is a non-steroidal antiinflammatory agent, inhibits COX activity, with IC_{59} S of 0.49 μ M and 36.6 μ M for COX-2 and COX-1, respectively.	O, O S'N OH O N	Meloxicam-13C,d3 is deuterium labeled Meloxicam. Meloxicam is a non-steroidal antiinflammatory agent, inhibits COX activity, with IC50s of 0.49 µM and 36.6 µM for COX-2 and COX-1, respectively.	о, о ^D D S N C D OH O N
Purity: 99.88% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 500 mg	inan in second	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	90 - 90 - 90 - 90 - 90 - 90 - 90 - 90 -
Malaviran d2		Malauiaam d2 1	
Meloxicam-03	Cat No HV-B02615	Meloxicam-d3-1	
Meloxicam-d3 is deuterium labeled Meloxicam. Meloxicam is a non-steroidal antiinflammatory agent, inhibits COX activity, with IC ₅₀ s of 0.49 μ M and 36.6 μ M for COX-2 and COX-1, respectively.		Meloxicam-d3-1 is the deuterium labeled Meloxicam. Meloxicam is a non-steroidal antiinflammatory agent, inhibits COX activity, with IC _{so} s of 0.49 μ M and 36.6 μ M for COX-2 and COX-1, respectively.	$(\begin{array}{c} \begin{array}{c} 0 \\ 0 \\ 0 \\ 0 \\ \end{array} \\ \end{array} \\ \begin{array}{c} 0 \\ 0 \\ \end{array} \\ \end{array} \\ \begin{array}{c} 0 \\ 0 \\ \end{array} \\ \begin{array}{c} 0 \\ 0 \\ \end{array} \\ \begin{array}{c} 0 \\ 0 \\ \end{array} \\ \end{array} \\ \begin{array}{c} 0 \\ 0 \\ \end{array} \\ \begin{array}{c} 0 \\ 0 \\ \end{array} \\ \end{array} \\ \begin{array}{c} 0 \\ 0 \\ \end{array} \\ \end{array} \\ \begin{array}{c} 0 \\ 0 \\ \end{array} \\ \end{array} \\ \begin{array}{c} 0 \\ 0 \\ \end{array} \\ \end{array} \\ \begin{array}{c} 0 \\ 0 \\ \end{array} \\ \end{array} \\ \begin{array}{c} 0 \\ 0 \\ \end{array} \\ \end{array} \\ \end{array} \\ \begin{array}{c} 0 \\ 0 \\ \end{array} \\ \end{array} \\ \end{array} \\ \begin{array}{c} 0 \\ 0 \\ \end{array} \\ \end{array} \\ \end{array} \\ \end{array} \\ \begin{array}{c} 0 \\ 0 \\ \end{array} \\$
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg, 10 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
Menin-MLL inhibitor MI-2	Cat. No.: HY-15222	Mensacarcin	Cat. No.: HY-122534
Menin-MLL inhibitor MI-2 is a $Menin-MLL$ interaction inhibitor with $\mathrm{IC}_{\mathrm{50}}$ of 446±28 nM.		Mensacarcin, a highly complex polyketide, strongly inhibits cell growth universally in cancer cell lines and potently induces apoptosis in melanoma cells.	
Purity:99.16%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg,	\$N 200 mg	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	о он





MG-277		MGCD-265 analog	
	Cat. No.: HY-130122		Cat. No.: HY-10991
MG-277, a molecular glue degrader, effectively induces degradation of a translation termination factor based on Cereblon E3 ligand, GSPT1, with a DC_{s0} of 1.3 nM.		MGCD-265 analog is a potent and oral active inhibitor of c-Met and VEGFR2 tyrosine kinases, with $IC_{so}s$ of 29 nM and 10 nM, respectively. MGCD-265 analog has significant antitumor activity.	artag
Purity:98.94%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg	a'	Purity: 98.57% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 50 mg	
MGH-CP1	Cat No : HV-139330	MI-1061	Cat. No : HY-125858
MGH-CP1 is a potent and orally active TEAD2 and TEAD4 auto-palmitoylation inhibitor with IC _{s0} s of 710 nM and 672 nM, respectively. MGH-CP1 can decrease the palmitoylation levels of endogenous or ectopically expressed TEAD proteins in cells.		MI-1061 is a potent, orally bioavailable, and chemically stable MDM2 (MDM2-p53 interaction) inhibitor (IC_{so} =4.4 nM; K_i =0.16 nM). MI-1061 potently activates p53 and induces apoptosis in the SJSA-1 xenograft tumor tissue in mice. Anti-tumor activity.	
Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Purity: 99.62% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg	
MI-1061 TFA		MI-136	Cat. No. 11/ 10210
$\begin{array}{ll} \mbox{MI-1061 TFA is a potent, orally bioavailable, and} \\ \mbox{chemically stable MDM2} (MDM2-p53 interaction) \\ \mbox{inhibitor} (IC_{so}=4.4 \mbox{ nN}; \mbox{K}_i=0.16 \mbox{ nM}). \mbox{MI-1061 TFA} \\ \mbox{potently activates } p53 \mbox{ and induces } apoptosis \mbox{ in} \\ \mbox{the SJSA-1 xenograft tumor tissue in mice.} \\ \mbox{Anti-tumor activity.} \\ \mbox{Purity: } 95.08\% \\ \mbox{Clinical Data: } No \mbox{ Development Reported} \\ \mbox{Size: } 1 \mbox{ mg, 5 \mbox{ mg, 10 \mbox{ mg}} \end{array}$		$\label{eq:memory_state} \begin{array}{ll} \mbox{MI-136 is an inhibitor of the menin-MLL} \\ \mbox{protein-protein interaction (PPI), with an IC_{so} of \\ 31 nM and a K_d of 23.6 nM. MI-136 shows to block \\ \mbox{AR signaling and has the potential for the study} \\ \mbox{in castration-resistant tumors.} \end{array}$	
MI-192	Cat. No. : HY-110264	MI-3 (Menin-MLL inhibitor 3)	Cat. No.: HY-15223
MI-192 is a selective HDAC2 and HDAC3 inhibitor with IC ₅₀ s of 30 nM and 16 nM, respectively. MI-192 is more selective for HDAC2/3 than other HDAC isomers.MI-192 induces myeloid leukaemic cells apoptosis . Anticaner and neuroprotective activities. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		$\label{eq:model} \begin{array}{ll} \text{MI-3} (\text{Menin-MLL inhibitor 3}) \text{ is a potent and high} \\ \text{affinity menin-MLL inhibitor with an IC}_{s0} \text{ of} \\ \text{648 nM and a } K_{d} \text{ of } 201 \text{ nM}. \\ \end{array}$	
Milademetan (DS-3032)	Cat. No. : HY-101266	Milademetan tosylate hydrate (DS-3032b; DS-3032 tosylate hydrate)	Cat. No.: HY-101266B
Milademetan (DS-3032) is a specific and orally active MDM2 inhibitor for the research of acute myeloid leukemia (AML) or solid tumors. Milademetan (DS-3032) induces G1 cell cycle arrest , senescence and apoptosis .	H'N C H H C H	Milademetan (DS-3032) tosylate hydrate is a specific and orally active MDM2 inhibitor for the research of acute myeloid leukemia (AML) or solid tumors. Milademetan (DS-3032) tosylate hydrate induces G1 cell cycle arrest, senescence and apoptosis.	and the state of t
Purity: 98.33% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg		Purity: 98.21% Clinical Data: Phase 2 Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg	н-н <u>Г</u>



MitoBloCK-6		MitoEbselen-2 chloride	
	Cat. No.: HY-122652	(MitoPeroxidase 2)	Cat. No.: HY-139381
MitoBloCK-6 is a potent Erv1/ALR inhibitor, with an IC _{s0} of 900 nM and 700 nM, respectively. MitoBloCK-6 also inhibits Erv2 (IC _{s0} =1.4 μ M). MitoBloCK-6 can induce apoptosis via cytochrome c release in hESCs. Purity: >98% Clinical Data: No Development Reported	CI CH NON	MitoEbselen-2 chloride (MitoPeroxidase 2), a mitochondria-targeted mimic of glutathione peroxidase, is a radiation mitigator. MitoEbselen-2 chloride is effective in reducing lipid hydroperoxides, preventing apoptotic cell death. Purity: >98% Clinical Data: No Development Reported	93.001 30
Size: 1 mg, 5 mg		Size: 1 mg, 5 mg	
Mitoguazone (Methylglyoxal-bis(guanylhydrazone); MGBG; Methyl-GAG)	Cat. No. : HY-106634	MitoTam bromide, hydrobromide	Cat. No.: HY-126222
Mitoguazone (Methylglyoxal-bis(guanylhydrazone)) is a synthetic polycarbonyl derivative with potent antineoplastic activity.	H2N HNNS NH	MitoTam bromide, hydrobromide, a Tamoxifen derivative, is an electron transport chain (ETC) inhibitor. MitoTam bromide, hydrobromide reduces mitochondrial membrane potential in senescent cells and affects mitochondrial morphology.	070
Purity:99.38%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg		Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	e de
MitoTam jodide, hydriedide		Mitotano	
Mitoran Iodide, nydnodide	Cat. No.: HY-126222A	(2,4'-DDD; o,p'-DDD)	Cat. No.: HY-13690
MitoTam iodide, hydriodide is a Tamoxifen derivative, an electron transport chain (ETC) inhibitor, spreduces mitochondrial membrane potential in senescent cells and affects mitochondrial morphology.		Mitotane(2,4'-DDD), an isomer of DDD and derivative of DDT, is an antineoplastic medication used in the treatment of adrenocortical carcinoma.	CI
Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg	¥	Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 500 mg	~ Ci
Mitotane-d4		Mivebresib	
(2,4'-DDD-d4; o,p'-DDD-d4)	Cat. No.: HY-13690S	(ABBV-075)	Cat. No.: HY-100015
Mitotane-d4 (2,4'-DDD-d4) is the deuterium labeled Mitotane. Mitotane (2,4'-DDD), an isomer of DDD and derivative of DDT, is an antineoplastic medication used in the treatment of adrenocortical carcinoma.		Mivebresib (ABBV-075) is a potent and orally active bromodomain and extraterminal domain (BET) bromodomain inhibitor. Mivebresib binds to BRD4 with a K _i of 1.5 nM.	
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity: 99.42% Clinical Data: Phase 1 Size: 10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 25 mg, 50	mg, 100 mg
MK-2206	Cat. No. : HY-108232	MK-2206 dihydrochloride (MK-2206 (2HCl))	Cat. No.: HY-10358
MK-2206 is an orally active, highly potent and selective allosteric Akt inhibitor, with IC_{50} of 8, 12, and 65 nM for Akt1, Akt2, and Akt3, respectively. Many breast cancer cell lines, and PIK3CA-mutant and cell lines with PTEN loss are sensitive to MK-2206. Anticancer activities.		MK-2206 dihydrochloride (MK-2206 (2HCl)) is an orally active allosteric AKT inhibitor with IC ₅₀ s of 5 nM, 12 nM, and 65 nM for AKT1 , AKT2 , and AKT3 , respectively. MK-2206 dihydrochloride induces autophagy .	
Clinical Data: No Development Reported Size: 1 mg, 5 mg		Clinical Data: Phase 2 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 10	200 mg

MK-4101		MK-8745	
MK-4101 is a Smoothened (SMO) antagonist (IC_{s0} of 1.1 μ M for 293 cells) and also a potent inhibitor of the hedgehog pathway (IC_{s0} of 1.5 μ M for mouse cells; IC_{s0} of 1 μ M for KYSE180	Cat. No.: HY-100036	MK-8745 is an $aurora\ A$ kinase inhibitor with an IC_{s0} of 0.6 nM.	Cat. No.: HY-13819
oesophageal cancer cells). Purity: 98.31% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 25 mg, 50	ng, 100 mg	Purity:99.49%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg, 50 mg, 100 mg	CI N H S
MK-886		MI 141	
(L 663536)	Cat. No.: HY-14166	(CID-2950007)	Cat. No.: HY-12755
MK-886 (L 663536) is a potent, cell-permeable and orally active FLAP (IC ₅₀ of 30 nM) and leukotriene biosynthesis (IC ₅₀ of 3 nM and 1.1 μ M in intact leukocytes and human whole blood, respectively) inhibitor. MK-886 is also a non-competitive PPAR α antagonist and can induce apoptosis . Purity: 99.74%	X S C C C C C C C C C C C C C C C C C C	ML141 (CID-2950007) is a potent, allosteric, selective and reversible non-competitive inhibitor of Cdc42 GTPase. ML141 inhibits Cdc42 wild type and Cdc42 Q61L mutant with EC_{50} s of 2.1 and 2.6 μ M, respectively. Purity: 99.71%	
Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	.00 mg	Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
ML291	Cat No. HV-101991	MMP2-IN-1	Cat No: HV-146754
ML291 is a UPR (unfolded protein response)-inducing sulfonamidebenzamide. ML291 overwhelms the adaptive capacity of the UPR and induces apoptosis in a variety of solid cancer models.		MMP2-IN-1 is a moderate potenet MMP2 inhibitor with IC_{s0} of 6.8 μ M. MMP2-IN-1 exhibits remarkable antiproliferative activity in certain cancer cells by arresting the cell cycle and inducing apoptosis .	
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	
MMPSI		MN58b	
	Cat. No.: HY-103346		Cat. No.: HY-108431
MMPSI is a potent and selective small molecule caspase 3 and caspase 7 inhibitor with an IC_{s_0} of 1.7 μ M for human caspase-3.	O, CH O	MN58b is a selective choline kinase α (CHKα) inhibitor, and results in inhibition of phosphocholine synthesis. MN58b reduces cell growth through the induction of apoptosis, and also has antitumoral activity.	1010.000'
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:99.17%Clinical Data:Size:1 mg	
MNK1/2-IN-6	Cat. No.: HY-146735	Mocetinostat (MGCD0103)	Cat. No.: HY-12164
MNK1/2-IN-6 is a potent and selective MNK1/2 inhibitor with IC_{so} s of 2.3 nM and 3.4 nM for MNK1 and MNK2, respectively. MNK1/2-IN-6 induces apoptosis in a concentration-dependent manner.	HAR CASO CAN	Mocetinostat (MGCD0103) is a potent, orally active and isotype-selective HDAC (Class I/IV) inhibitor with IC ₅₀ s of 0.15, 0.29, 1.66 and 0.59 μ M for HDAC1, HDAC2, HDAC3 and HDAC11, respectively. Mocetinostat shows no inhibition on HDAC4, HDAC5, HDAC6, HDAC7, or HDAC8.	and the
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity: 99.43% Clinical Data: Phase 2 Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg	

Moexipril hydrochloride		Moexipril-d5	
(RS-10085)	Cat. No.: HY-B0378A		Cat. No.: HY-117281S
Moexipril hydrochloride is a potent orally active non-sulfhydryl angiotensin converting enzyme (ACE) inhibitor, which is used for the treatment of hypertension and congestive heart failure.	Contraction of the second seco	Moexipril-d5 is the deuterium labeled Moexipril. Moexipril hydrochloride is a potent orally active non-sulfhydryl angiotensin converting enzyme(ACE) inhibitor, which is used for the treatment of hypertension and congestive heart failure.	
Purity: 98.95% Clinical Data: Launched Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg	HCI	Purity:>98%Clinical Data:Size:1 mg, 10 mg	bi dese
Moexipril-d5 hydrochloride	Cat. No.: HY-B0378AS	Momelotinib (CYT387)	Cat. No.: HY-10961
Moexipril-d5 (hydrochloride) is deuterium labeled Moexipril (hydrochloride).		Momelotinib (CYT387) is an ATP-competitive inhibitor of JAK1/JAK2 with $IC_{50}a$ of 11 nM and 18 nM,respectively. CYT387 shows much less activity against JAK3.	ooboh.
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	b	Purity: 98.93% Clinical Data: Phase 3 Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	200 mg
Momelotinib sulfate	Cat. No. UV 10062	Momordin Ic	Cot No. UV N0220
Momelotinib sulfate (CYT387 sulfate salt) is an ATP-competitive inhibitor of JAK1/JAK2 with IC_{50} of 11 nM/18 nM, 10-fold selectivity versus JAK3 (IC_{50} =155 nM).		Momordin Ic is a principal saponin constituent of Fructus Kochiae, with with anti-cancer bioactivity. Momordin Ic induces apoptosis through oxidative stress-regulated mitochondrial dysfunction. Purity: 99.71% Clinical Data: No Development Reported	
Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Size: 5 mg, 10 mg, 20 mg	
Monastrol ((±)-Monastrol)	Cat. No.: HY-101071A	Monensin	Cat. No.: HY-N4302
Monastrol is a potent and cell-permeable inhibitor of the mitotic kinesin Eg5 with an IC _{so} value of 14 μM. Purity: 99.70% Clinical Data: No Development Reported	HN + O S + N + O H	Monensin is a naturally occurring bioactive ionophore produced by Streptomyces spp. Monensin can bind protons and monovalent cations. Monensin exhibits a broad spectrum activity against opportunistic pathogens of humans in both drug sensitive and resistant strains. Purity: >98% Clinical Data: No Development Reported	PH OF OF OF
Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 25 mg, 50) mg, 100 mg	Size: 5 mg	
Monomethyl auristatin E (MMAE; SGD-1010; Vedotin)	Cat. No. : HY-15162	Morroniside	Cat. No.: HY-N0532
Monomethyl auristatin E (MMAE; SGD-1010) is a synthetic derivative of dolastatin 10 and functions as a potent mitotic inhibitor by inhibiting tubulin polymerization.		Morroniside has neuroprotective effect by inhibiting neuron apoptosis and MMP2/9 expression.	
Purity: 99.92% Clinical Data: Phase 2 Size: 5 mg, 10 mg, 50 mg, 100 mg, 500 mg, 1 g	0	Purity:98.55%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 25 mg	

MPI-0479605	Cat No : HY-12660	MPP hydrochloride	Cat No : HY-103454B
MPI-0479605 is a potent and selective ATP-competitive inhibitor of Mps1 , with an IC _{so} of 1.8 nM.		MPP hydrochloride is a potent and selective ER (estrogen receptor) modulator. MPP hydrochloride induces significant apoptosis in the endometrial cancer and oLE cell lines. MPP hydrochloride reverses the positive effects of beta-estradiol.	HO CH-CO-CH-CO-CH H-CI
Clinical Data: No Development Reported Size: 10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 50 mg		Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
MPT0B392	Cat. No.: HY-101287	MPT0E028	Cat. No. : HY-124295
MPT0B392, an orally active quinoline derivative, induces c-Jun N-terminal kinase (JNK) activation, leading to apoptosis .	NH2 N Store	MPT0E028 is an orally active and selective HDAC inhibitor with IC_{50} s of 53.0 nM, 106.2 nM, 29.5 nM for HDAC1, HDAC2 and HDAC6, respectively.	HO, H
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	~	Purity:>98%Clinical Data:Phase 1Size:1 mg, 5 mg	o
MPTP hydrochloride	Cat. No. : HY-15608	MRIA9	Cat. No. : HY-139253
MPTP hydrochloride is a brain penetrant dopamine neurotoxin, inducing Parkinson's Disease. MPTP hydrochloride, a precusor of MPP+, induces apoptosis .		MRIA9 is an ATP-competitive, pan Salt-Inducible kinase (SIK) and PAK2/3 inhibitor, with IC_{s0} values of 516 nM, 180 nM and 127 nM for SIK1, SIK2 and SIK3, respectively.	
Purity:99.54%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg, 50 mg, 100 mg	H-CI	Purity:98.10%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg	0. NH2
MRS 2578	Cat. No. : HY-13104	MRT199665	Cat. No. : HY-120877
MRS 2578 is a selective and potent P2Y6 receptor antagonist with IC_{so} s of 37 nM (human) and 98 nM (rat). MRS 2578 exhibits insignificant activity at P2Y1, P2Y2, P2Y4, and P2Y11 receptors.	~ ^{c⁴O²}³²~³²s^Ov^{c⁴}}	MRT199665 is a potent and ATP-competitive, selective MARK/SIK/AMPK inhibitor with IC ₅₀ s of 2/2/3/2 nM, 10/10 nM, and 110/12/43 nM for MARK1/MARK2/MARK3/MARK14, AMPKα1/AMPKα2, and SIK1/SIK2/SIK3, respectively.	ano toto
Purity:98.15%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg, 50 mg		Purity: 99.73% Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg, 50 mg	4.54
MS1943	Cat. No.: HY-133129	MSN-125	Cat. No.: HY-120079
MS1943 is a first-in-class, orally bioavailable EZH2 selective degrader, with an IC_{s0} of 120 nM. MS1943 significantly reduces EZH2 protein levels in numerous triple-negative breast cancer (TNBC) and other cancer and noncancerous cell lines.	the Photosia	MSN-125 is a potent Bax and Bak oligomerization inhibitor. MSN-125 prevents mitochondrial outer membrane permeabilization (MOMP) with an IC ₅₀ of 4 μ M.	
Purity:98.18%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg		Purity:98.64%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg	

MT 63-78	Cat. No.: HY-W058849	mTOR/HDAC6-IN-1	Cat. No.: HY-144449
MT 63-78 is a specific and potent direct AMPK activator with an EC _{s0} of 25 µM. MT 63–78 also induces cell mitotic arrest and apoptosis . MT 63-78 blocks prostate cancer growth by inhibiting the lipogenesis and mTORC1 pathways. MT 63-78 has antitumor effects. Purity: 98.22% Clinical Data: No Development Reported Size: 5 mg, 10 mg	CHOH OH CH ZH	mTOR/HDAC6-IN-1 is a potent mTOR and HDAC6 dual inhibitor (IC ₅₀ S of 133.7 nM and 56 nM for mTOR and HDAC6, respectively). mTOR/HDAC6-IN-1 can induce significant autophagy , apoptosis and suppress migration. mTOR/HDAC6-IN-1 has potential to research Triple-negative breast cancer (TNBC). Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	CI C
Multi-kinase-IN-1	Cat. No.: HY-146014	MV1	Cat. No. : HY-113534
Multi-kinase-IN-1 (Compound 11k) is a potent kinase inhibitor with antitumor activity. Multi-kinase-IN-1 induces cell apoptosis , and can be studied for colorectal cancer .	Str. C.	MV1 is an antagonist of IAP (inhibitor of apoptosis protein), leads to protein knockdown of HaloTag-fused proteins when combined with HaloTag ligand. Purity: 99.54% Clinical Data: No Development Reported	
Size: 1 mg, 5 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
MYCMI-6 (NSC354961)	Cat. No.: HY-124675	Mycophenolate Mofetil (RS 61443; TM-MMF)	Cat. No.: HY-B0199
MYCMI-6 (NSC354961) is a potent and selective endogenous MYC:MAX protein interactions inhibitor. MYCMI-6 blocks MYC-driven transcription and binds selectively to the MYC bHLHZip domain with a K_d of 1.6µM.	NN (NN) (N	Mycophenolate mofetil (RS 61443) is the morpholinoethylester prodrug of Mycophenolic acid. Mycophenolate mofetil inhibits de novo purine synthesis via the inhibition of inosine monophosphate dehydrogenase (IMPDH) .	it is a compared to the second
Purity:95.95%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 50 mg, 100 mg		Purity: 99.68% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 200 mg, 1 g, 5 g	
Mycophenolate Mofetil-d4	Cat. No. : HY-B0199S	Mycophenolic acid (Mycophenolate)	Cat. No.: HY-B0421
Mycophenolate Mofetil-d4 is the deuterium labeled Mycophenolate Mofetil. Mycophenolate mofetil (RS 61443) is the morpholinoethylester prodrug of Mycophenolic acid.	Jan and a star	Mycophenolic acid is a potent uncompetitive inosine monophosphate dehydrogenase (IMPDH) inhibitor with an EC _{so} of 0.24 µM. Mycophenolic acid demonstrates antiviral effects against a wide range of RNA viruses including influenza	offor offor
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 50 mg		Purity:99.87%Clinical Data:LaunchedSize:10 mM × 1 mL, 100 mg, 500 mg, 1 g	
Mycophenolic acid 13C,D3 (Mycophenolate 13C,D3)	Cat. No. : HY-B0421S1	Myricetin (Cannabiscetin)	Cat. No.: HY-15097
Mycophenolic acid 13C,D3 (Mycophenolate 13C,D3) is deuterium labeled Mycophenolic acid 13C. Mycophenolic acid is an an immunosuppresant drug and has potent anti-proliferative activity.	ано с он	Myricetin is a common plant-derived flavonoid with a wide range of activities including strong anti-oxidant, anticancer, antidiabetic and anti-inflammatory activities.	но он он он
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity: 98.08% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 50 mg, 100 mg, 200 mg, 500 mg	un u

Myristoleic acid		Mytoxin B	
	Cat. No.: HY-113332		Cat. No.: HY-131055
Myristoleic acid, a cytotoxic component in the		Mytoxin B is an ADC cytotoxin. Mytoxin B is a	7°
extract from Serenoa repens, induces apoptosis and necrosis in human prostatic LNCaP cells		satratoxin-type trichothecene macrolide and is similar to the effect of LY294002 (HY-10108)	9R
	l	Mytoxin B induces cell apoptosis via PI3K/Akt	но
	~~~~~~~ 0H	pathway.	
Duritar > 00.00/		Durit restance (1997)	
Clinical Data: No Development Reported		Clinical Data: No Development Reported	Ho
Size: 100 mg		Size: 1 mg, 5 mg	
n-Octyl caffeate		NAF-IN-M22	
	Cat. No.: HY-N8398		Cat. No.: HY-115537
n-Octyl catteate shows anti-cancer and apoptosis inducing activity in highly liver-metastatic		NAE-IN-M22 is a potent, selective and reversible inhibitor of NEDD8 activating enzyme (NAE) with	
murine colon 26-L5 carcinoma cell lines.	<u></u>	potency in micromolar range. NAE-IN-M22 inhibits	
	Harrison	multiple cancer cell lines and induces apoptosis	TT, OTO
	HO	in A549 cells. NAE-IN-M22 also can inhibit tumor growth in vivo.	н
Purity: >98%		Purity: 99.67%	
Clinical Data: No Development Reported		Clinical Data: No Development Reported	
Size: 5 mg, 10 mg, 25 mg		Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
Nafamostat		Nafamostat hydrochloride	
	Cat. No.: HY-B0190		Cat. No.: HY-B0190B
Nafamostat, a synthetic serine protease inhibitor.		Nafamostat hydrochloride, a synthetic serine	
is an anticoagulant. Nafamostat supresses T cell		protease inhibitor, is an anticoagulant.	
auto-reactivity by decreasing granzyme activity	^R _{NH₂}	Nafamostat hydrochloride supresses T cell	
SARS-CoV-2.	HALLOOT S MH	and CTL cytolysis. Nafamostat hydrochloride blocks	HALLI I
	NH	activation of SARS-CoV-2.	HCI HCI
Purity: >98%		Purity: >98%	
Clinical Data: Launched		Clinical Data: Launched	
Size. I mg, 5 mg		<b>Size.</b> 1 mg, 5 mg	
Nafamostat mesylate		Nampt-IN-3	
(FUT-175)	Cat. No.: HY-B0190A		Cat. No.: HY-108701
Nafamostat mesylate, a synthetic serine protease		Nampt-IN-3 (Compound 35) simultaneously inhibit	
inhibitor, is an anticoagulant. Nafamostat	, R_NH,	nicotinamide phosphoribosyltransferase (NAMPT) and HDAC with IC is of 31 pM and 55 pM	ml
decreasing granzyme activity and CTL cytolysis.	HALOO B	respectively. Nampt-IN-3 effectively induces cell	
Nafamostat mesylate blocks activation of	NH 0 0 	apoptosis and autophagy and ultimately leads to	S-m-m
SARS-CoV-2.	0 0	cell death.	$\langle \mathcal{Q} \rangle$
Purity: 98.06%		Purity: 99.27%	
Size: 10 mM × 1 mL, 10 mg, 50 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	)0 mg
Nanhthazarin			
(DHNO: 5.8-Dibydroxy-1.4-paphthoguinopo)	Cat No : HV N7526	NDDHEA	Cat No + HV 125219
	Cat. NO., FIT-IN/320		Cat. INO., FIT-100016
Naphthazarin (DHNQ) is a naturally occurring	он о	NBDHEX is a potent glutathione S-transferase P1-1	
compound.	$\checkmark$	tumor cells.	0
	ſ Ĩ Ì		0 ^{,N*}
	$\searrow$		N° Y S
Purity: 9813%		Purity: 98 58%	
Clinical Data: No Development Reported	UH U	Clinical Data: No Development Reported	
Size: 100 mg, 250 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	

NCT-58	<b>Cat. No.</b> : HY-145102	NCX4040 (NO-Aspirin)	<b>Cat. No.:</b> HY-103385
NCT-58 is a potent inhibitor of C-terminal <b>HSP90</b> . <b>Purity:</b> >98% <b>Clinical Data:</b> No Development Reported		NCX4040 (NO-Aspirin), a non-steroidal anti-inflammatory drug (NSAID), is a nitric oxide (NO) releasing form of Aspirin. NCX4040 induces <b>apoptosis</b> in PC3 metastatic prostate cancer cells. NCX4040 has anti-inflammatory and anti-cancer effects. <b>Purity:</b> >98% <b>Clinical Data:</b> No Development Reported	
Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg		Size: 1 mg, 5 mg	
Nebivolol hydrochloride		Necrostatin-7	
(R 065824 hydrochloride)	Cat. No.: HY-B0203A	(Nec-7)	Cat. No.: HY-117200
Nebivolol hydrochloride selectively inhibits $\beta$ 1- adrenergic receptor with IC50 of 0.8 nM. Target: $\beta$ 1- adrenergic receptor Nebivolol reduces cell proliferation of human coronary smooth muscle cells (haCSMCs) and endothelial cells (haECs) in a concentration- and time-dependent maner.	FUTURE HOLE FOR	Necrotatin-7 (Nec-7) is a potent <b>necroptosis</b> inhibitor with an $EC_{so}$ of 10.6 $\mu$ M. Necrotatin-7 does not inhibit recombinant RIP1 kinase.	
Clinical Data: Launched		Clinical Data: No Development Reported	
Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 500 mg		Size: 1 mg, 5 mg	
Neferine		Nelarabine	
((-)-Neferine)	Cat. No.: HY-N0441	(506U78; GW 506U78; Nelzarabine)	Cat. No.: HY-13701
Neferine is a major bisbenzylisoquinline alkaloid. Neferine strongly inhibits <b>NF-κB</b> activation.	dag. or	Nelarabine (Arranon, 506U78) is a purine nucleoside analog and DNA synthesis inhibitor with IC50 from 0.067-2.15 $\mu$ M in tumor cells. Nelarabine is a chemotherapy drug used in T-cell acute lymphoblastic leukemia.	
Purity:         99.92%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg		Purity:         99.88%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
Nemorosone		Neobavaisoflavone	
Nemorosone	Cat. No.: HY-121458	Neobavaisonavone	Cat. No.: HY-N0720
Nemorosone is the main component of the floral resin of Clusia rosea. Nemorosone has an antiproliferative effect on cancer cells. Nemorosone induces <b>apoptosis</b> in HT-29 and LoVo cells.	Horo	Neobavaisoflavone, a flavonoid, is isolated from the seeds of Psoralea corylifolia. Neobavaisoflavone exhibits anti-inflammatory, anti-cancer and anti-oxidation activities. Neobavaisoflavone inhibits <b>DNA polymerase</b> at moderate to high concentrations.	HO COLON
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg	~	Purity:99.91%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg	
Neocarzinostatin	<b>Cat. No.:</b> HY-111183	Neoechinulin A	<b>Cat. No.:</b> HY-N3204
Neocarzinostatin, a potent <b>DNA-damaging</b> , anti-tumor antibiotic, recognizes double-stranded DNA bulge and induces DNA double strand breaks ( <b>DSBs</b> ). Neocarzinostatin induces <b>apoptosis</b> . Neocarzinostatin has potential for EpCAM-positive cancers treatment.	Neocarzinostatin	Neoechinulin A is an isoprenyl indole alkaloid that exhibits scavenging, neurotrophic factor-like, and anti-apoptotic activities. Neoechinulin A induces memory improvements and antidepressant-like effects in mice.	
Purity:     ≥90.0%       Clinical Data:     No Development Reported       Size:     100 μg		Purity:     ≥98.0%       Clinical Data:     No Development Reported       Size:     1 mg	∕ N H

Neogambogic acid		Neoxanthin	
Neogambogic acid, an active ingredient in garcinia, induces apoptosis and has anticancer effect. Neogambogic acid has significant inhibitory activity toward methicillin-resistant Staphylococcus aureus (MRSA). Purity: >98% Clinical Data: No Development Reported Size: 5 mg, 10 mg, 20 mg		Neoxanthin is a major xanthophyll carotenoid and a precursor of the plant hormone abscisic acid in dark green leafy vegetables. Neoxanthin is a potent antioxidant and light-harvesting pigment. Neoxanthin induces <b>apoptosis</b> and has anticancer actions.         Purity:       ≥99.0%         Clinical Data:       No Development Reported         Size:       1 mg	Ha Carl NO. HT-IN/323
<mark>Neriifolin</mark> (17β-Neriifolin)	<b>Cat. No</b> .: HY-N8441	Nerol	<b>Cat. No.</b> : HY-N7063
Neriifolin, a CNS-penetrating cardiac glycoside, is an inhibitor of the Na ⁺ , K ⁺ -ATPase. Neriifolin can target beclin 1, inhibits the formation of LC3-associated phagosomes and ameliorates experimental autoimmune encephalomyelitis (EAE) development.	$H_{0} \xrightarrow{f_{0}}_{1} O_{0} O_{$	Nerol is a constituent of neroli oil. Nerol Nerol triggers mitochondrial dysfunction and induces apoptosis via elevation of Ca ²⁺ and <b>ROS</b> . <b>Antifungal</b> activity.	HO
Purity:     ≥96.0%       Clinical Data:     No Development Reported       Size:     5 mg		Purity:         ≥97.0%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 10 mg, 50 mg, 100 mg	
Nevanimibe	Cat. No. 11V 100200	Nevanimibe hydrochloride	
(PD-132301; AIR-101)Nevanimibe (PD-132301) is an orally active and selective acyl-coenzyme A:cholesterol O-acyltransferase 1 (ACAT1) inhibitor with an $EC_{s0}$ of 9 nM. Nevanimibe inhibits ACAT2 with an $EC_{s0}$ of 368 nM. Nevanimibe induces cell apoptosis and has the potential for adrenocortical cancer.Purity:>98% Clinical Data:Phase 2 Size:1 mg, 5 mg		(PD-132301 hydrochloride; ATR101 hydrochloride)Nevanimibe hydrochloride (PD-132301 hydrochloride)is an orally active and selective acyl-coenzymeA:cholesterol O-acyltransferase 1 (ACAT1)inhibitor with an EC ₅₀ of 9 nM. Nevanimibehydrochloride inhibits ACAT2 with an EC ₅₀ of 368nM.Purity:98.07%Clinical Data:Phase 2Size:10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	HN HY-100399A
NF-ĸB-IN-4	<b>Cat. No.:</b> HY-144765	NHWD-870	<b>Cat. No.</b> : HY-134463
NF-κB-IN-4 (compound 17) is a potent and BBB-penetrated NF-κB pathway inhibitor with blood brain barrier (BBB) permeability. NF-κB-IN-4 exhibits potential anti-neuroinflammatory activity with low toxicity. Purity: >98% Clinical Data: No Development Reported	NH2 N N F	NHWD-870 is a potent, orally active and selective         BET family bromodomain inhibitor and only         binds bromodomains of BRD2, BRD3, BRD4 (IC ₅₀ =2.7         nM), and BRDT. NHWD-870 has potent tumor         suppressive efficacies and suppresses cancer         cell-macrophage interaction.         Purity:       99.36%         Clinical Data:       No Development Reported	
Size: 1 mg, 5 mg		Size:         5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
Nigericin	<b>Cat. No.</b> : HY-127019	Nimbolide	<b>Cat. No.:</b> HY-116035
Nigericin is an <b>antibiotic</b> derived from Streptomyces hygroscopicus that act as a $K^{+}/H^{+}$ <b>ionophore</b> , promoting $K^{+}/H^{+}$ exchange across mitochondrial membranes.Nigericin can be a <b>NLRP3</b> activator that induces the release of IL-1 $\beta$ as a NALP3-dependent manner.		Nimbolide is a triterpene derived from the leaves and flowers of neem (Azadirachta indica L). Nimbolide induces apoptosis through inactivation of NF-KB. Nimbolide inhibits CDK4/CDK6 kinase activity. Nimbolide suppresses the NF-KB, Wnt, PI3K-Akt, MAPK and JAK-STAT signaling pathways.	
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg		Purity:     99.94%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	

Nimustine bydrochloride		Niranarih	
(ACNU)	Cat. No.: HY-13703A	(MK-4827)	<b>Cat. No.</b> : HY-10619
Nimustine hydrochloride (ACNU) is a DNA cross-linking and DNA alkylating agent, which induces DNA replication blocking lesions and DNA double-strand breaks and inhibits <b>DNA synthesis</b> , commonly used in chemotherapy for glioblastomas. <b>Purity:</b> 99.90% <b>Clinical Data:</b> Launched	H-CI	Niraparib (MK-4827) is a highly potent and orally bioavailable <b>PARP1</b> and <b>PARP2</b> inhibitor with IC ₅₀ S of 3.8 and 2.1 nM, respectively. Niraparib leads to inhibition of repair of DNA damage, activates <b>apoptosis</b> and shows anti-tumor activity. <b>Purity:</b> 99.97% <b>Clinical Data:</b> Launched	
Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
Niraparib hydrochloride (MK-4827 hydrochloride)	<b>Cat. No.:</b> HY-10619A	Niraparib tosylate (MK-4827 tosylate)	<b>Cat. No.:</b> HY-10619B
Niraparib hydrochloride (MK-4827 hydrochloride) is         a highly potent and orally bioavailable PARP1 and         PARP2 inhibitor with IC ₅₀ S of 3.8 and 2.1 nM,         respectively. Niraparib hydrochloride leads to         inhibition of repair of DNA damage, activates         apoptosis and shows anti-tumor activity.         Purity:       99.80%         Clinical Data:       Launched         Size:       10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		eq:space-	
Nirogacestat (PF-3084014; PF-03084014)	<b>Cat. No.</b> : HY-15185	Nirogacestat dihydrobromide (PF-3084014 dihydrobromide; PF-03084014 dihydrobromid	e) Cat. No.: HY-15185B
Nirogacestat (PF-3084014) is a reversible, orally bioavailable, noncompetitive, and selective $\gamma$ -secretase inhibitor with an IC _{s0} of 6.2 nM.	and the state of the	Nirogacestat dihydrobromide (PF-3084014 dihydrobromide) is a reversible, orally bioavailable, noncompetitive, and selective $\gamma$ -secretase inhibitor with an IC ₅₀ of 6.2 nM.	PLOT THE HER
Purity:         98.76%           Clinical Data:         Phase 3           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
Nitidine chloride	<b>Cat. No.:</b> HY-N0498	Nitroaspirin (NCX 4016)	<b>Cat. No.:</b> HY-123823
Nitidine chloride, a potential <b>anti-malarial</b> lead compound derived from Zanthoxylum nitidum (Roxb) DC, exerts potent anticancer activity through diverse pathways, including inducing <b>apoptosis</b> , inhibiting <b>STAT3</b> signaling cascade, <b>DNA topoisomerase 1 and 2A, ERK</b> and		Nitroaspirin (NCX 4016) is a nitric oxide (NO) donor and a nitro-derivative of Aspirin, which combines with Nitroaspirin to inhibit cyclooxygenase.	
Purity:       99.61%         Clinical Data:       No Development Reported         Size:       5 mg, 10 mg, 20 mg		Purity:       >98%         Clinical Data:       Phase 2         Size:       5 mg, 10 mg, 50 mg, 100 mg	
Nivalenol	<b>Cat. No.:</b> HY-N6801	NKP-1339 (IT-139; KP-1339)	<b>Cat. No.:</b> HY-16350
Nivalenol, classified as type B trichotecenes toxins produced by Fusarium graminearum, is a fungal metabolite present in agricultural product. Nivalenol induces cell death through <b>caspase</b> -dependent mechanisms and via the intrinsic <b>apoptotic</b> pathway.	OH OH OH	NKP-1339 (IT-139; KP-1339) is the first-in-class ruthenium-based anticancer agent in development against solid cancer with limited side effects. NKP-1339 induces G2/M cell cycle arrest, blockage of <b>DNA synthesis</b> , and induction of <b>apoptosis</b> via the mitochondrial pathway.	
Purity:     ≥99.0%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg		Purity:         98.14%           Clinical Data:         Phase 1           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	762 °

Nobiletin	<b>Cat. No.:</b> HY-N0155	Nocodazole (Oncodazole; R17934)	<b>Cat. No.:</b> HY-13520
Nobiletin is a poly-methoxylated flavone from the citrus peel that improves memory loss. Nobiletin is a <b>retinoid acid receptor-related orphan receptors (RORs)</b> agonist.		Nocodazole (Oncodazole) is a rapidly-reversible inhibitor of <b>microtubule</b> . Nocodazole binds to $\beta$ -tubulin and disrupts microtubule assembly/disassembly dynamics, which prevents mitosis and induces apoptosis in tumor cells.	STOLEN SO
Purity:         99.52%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	Pruda Inda	Purity:99.66%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg, 50 mg, 100 mg	
Nonactin (Ammonium ionophore I)	<b>Cat. No.:</b> HY-N6790	nor-NOHA acetate (Νω-Hydroxy-nor-L-arginine acetate)	<b>Cat. No.:</b> HY-112885A
Nonactin is a naturally occurring macrotetrolide antibiotic from Streptomyces griseus. Nonactin acts as an ionophore for monovalent cations, including K*, and NH ₄ *. Nonactin is able to uncouple the oxidative phosphorylation (OXPHOS) of mitochondria. Purity: $\geq$ 99.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 1 mg, 5 mg		nor-NOHA acetate (N $\omega$ -Hydroxy-nor-L-arginine acetate) is a specific and reversible <b>arginase</b> inhibitor, induces apoptosis in ARG2-expressing cells under hypoxia but not normoxia.Anti-leukemic activity, effective in endothelial dysfunction, immunosuppression and metabolism.Purity: $\geq$ 98.0%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg	но Н Н Рон NH NH2 Сон Сон
Nortrachelogenin		Nortriptyline-d3 hydrochloride (Desmethylamitrip	tyline-d3
((-)-Wikstromol; (-)-Nortrachelogenin) Nortrachelogenin ((-)-Wikstromol) from Partrinia scabiosaefolia elicits an apoptotic response in Candida albicans. Purity: >98%	Cat. No.: HY-N3171 HO $++++++++++++++++++++++++++++++++++++$	hydrochloride; Desitriptilina-d3 hydrochloride) Nortriptyline-d3 (Desmethylamitriptyline-d3) hydrochloride is the deuterium labeled Nortriptyline hydrochloride. Purity: >98%	Cat. No.: HY-B14175
Clinical Data: No Development Reported Size: 1 mg, 5 mg		Clinical Data:No Development ReportedSize:2.5 mg, 1 mg, 5 mg, 10 mg	
Notoginsenoside R1 (Sanchinoside R1; Sanqi glucoside R1)	<b>Cat. No.:</b> HY-N0615	Notopterol	<b>Cat. No.:</b> HY-N0564
Notoginsenoside R1 (Sanchinoside R1), a saponin, is isolated from P. notoginseng. Notoginsenoside R1 exhibits anti-oxidation, anti-inflammatory, anti-angiogenic, and anti-apoptosis activities.		Notopterol is a coumarin extracted from N. incisum. Notopterol induces <b>apoptosis</b> and has antipyretic, analgesic and anti-inflammatory effects. Notopterol is used for acute myeloid leukemia (AML). <b>Purity:</b> 99.27%	of of the
Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg		Clinical Data:       No Development Reported         Size:       5 mg, 10 mg, 20 mg	
NPS-1034	<b>Cat. No.:</b> HY-100509	NQDI-1	<b>Cat. No.:</b> HY-19566
NPS-1034 is a dual inhibitor of AXL and MET with $\mathrm{IC}_{50}$ s of 10.3 and 48 nM, respectively.	-o-jia	NQDI-1 inhibits apoptosis signal-regulating kinase 1 (ASK1) with a K, of 500 nM and an IC $_{\rm 50}$ of 3 $\mu M.$	O O NH
Purity:         ≥98.0%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	)0 mg	Purity:         95.93%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	U O

NS-1619	<b>C</b> + <b>N</b> + 12400	NS3694	C + N + 10 100250
NS-1619 is an opener of large conductance $Ca^{2*}$ -activated K ⁺ (BK) channel. NS-1619 is a highly effective relaxant with an EC ₅₀ of about 10–30µM in several smooth muscles of blood vessels and other tissues. Purity: $\geq$ 98.0% Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg	HO - FFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFFF	NS3694, a diarylurea compound, is an <b>apoptosome</b> inhibitor. NS3694 inhibits apoptosome formation and caspase activation. <b>Purity:</b> >98% <b>Clinical Data:</b> No Development Reported <b>Size:</b> 1 mg, 5 mg	
NSC 146109 hydrochloride	<b>Cat. No.:</b> HY-108638	NSC 15364	<b>Cat. No.</b> : HY-108937
NSC 146109 hydrochloride is a small-molecule <b>p53</b> activator that target <b>MDMX</b> and can be used for breast cancer research. NSC 146109 hydrochloride is a pseudourea derivative, promotes breast cancer cells to undergo <b>apoptosis</b> through activating p53 and inducing expression of proapoptotic genes. <b>Purity:</b> 99.60% <b>Clinical Data:</b> No Development Reported Since 5 mg 10 mg 25 mg 50 mg 100 mg	HN NH2 S S H-Cl	NSC 15364 is an inhibitor of VDAC1 oligomerization and apoptosis. Purity: 99.27% Clinical Data: No Development Reported Size 10 mM x 1 mJ 10 mg 50 mg 100 mg 250 mg	HAN TO BE AND
Size. 5 mg, 10 mg, 25 mg, 50 mg, 100 mg		Size. 10 milli × 1 mil, 10 mill, 30 mill, 100 mill, 250 mill	
NSC 23766 trihydrochloride	<b>Cat. No.:</b> HY-15723A	NSC 95397	<b>Cat. No.</b> : HY-108543
NSC 23766 trihydrochloride is an inhibitor of <b>Rac1</b> activation.	Ha Ha Ha	NSC 95397 is a potent, selective <b>Cdc25</b> dual specificity phosphatase inhibitor ( $K_i$ =32 nM (Cdc25A), 96 nM (Cdc25B), 40 nM (Cdc25C); <b>IC</b> ₅₀ =22.3 nM (human Cdc25A), 56.9 nM (human Cdc25C), 125 nM (Cdc25B)).	С С С С С С С С С С С С С С С С С С С
Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg		Purity:         98.02%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	0 mg
NSC-87877	<b>Cat. No.:</b> HY-18756	NSC-87877 disodium	<b>Cat. No.:</b> HY-18756A
NSC-87877 is a potent inhibitor of Shp2 and Shp1 protein tyrosine phosphatases (SH-PTP2 and SH-PTP1), with $IC_{50}$ values of 0.318 $\mu$ M, 0.355 $\mu$ M shp2 and shp1, respectively. NSC-87877 also inhibits dual-specificity phosphatase 26 (DUSP26).	HO. 0 O U N.N. UH.N. O = = = O OH	NSC-87877 disodium is a potent inhibitor of Shp2 and Shp1 protein tyrosine phosphatases (SH-PTP2 and SH-PTP1), with IC ₅₀ values of 0.318 $\mu$ M, 0.355 $\mu$ M shp2 and shp1, respectively. NSC-87877 also inhibits dual-specificity phosphatase 26 (DUSP26).	CNu 0-5-0 N CH N C C O O CNU
Purity:         ≥98.0%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	0 mg	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
NSC348884	<b>Cat. No.:</b> HY-13915	NSC697923	<b>Cat. No.:</b> HY-13811
NSC348884 is a nucleophosmin inhibitor disrupts oligomer formation and induces apoptosis, inhibits cell proliferation at an IC50 of 1.7-4.0 $\mu$ M in distinct cancer cell lines.		NSC697923 is a potent <b>UBE2N</b> (ubiquitin-conjugating enzyme E2 N, Ubc13) inhibitor. NSC697923 induces neuroblastoma (NB) cell death via promoting nuclear importation of p53 in p53 wild-type NB cells.	O'A O NO
Purity:         ≥98.0%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	0 mg	Purity:99.16%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	

NSC745885	<b>Cat. No.:</b> HY-119198	NTR 368	<b>Cat. No.</b> : HY-P1176
NSC745885 an effective <b>anti-tumor</b> agent, shows selective toxicity against multiple cancer cell lines but not normal cells. NSC745885 is an effective down-regulator of <b>EZH2</b> via proteasome-mediated degradation. <b>Purity:</b> ≥98.0% <b>Clinical Data:</b> No Development Reported <b>Size:</b> 5 mg, 10 mg, 50 mg, 100 mg	O N-S N O	NTR 368 is a peptide derived from p75 neurotrophin receptor (p75NTR) corresponding to residues 368-381 of the human receptor. NTR 368 has helix forming propensity in the presence of micellar lipid. NTR 368 is a potent inducer of neural apoptosis.Purity:>98%Clinical Data:No Development Reported Size:1mg, 5	Ac-ATLDALLAALRRIQ-NH2
NTR 368 TFA	<b>Cat. No.:</b> HY-P1176A	NU 7026 (LY293646)	<b>Cat. No.:</b> HY-15719
NTR 368 TFA is a peptide derived from p75 neurotrophin receptor (p75NTR) corresponding to residues 368-381 of the human receptor. NTR 368 TFA has helix forming propensity in the presence of micellar lipid. NTR 368 TFA is a potent inducer of neural <b>apoptosis</b> . <b>Purity:</b> >98%	Ac-ATLDALLAALRRID-NH2 (TFA IsH)	NU 7026 (LY293646) is a novel specific <b>DNA-PK</b> inhibitor with IC _{so} of 0.23 $\mu$ M, also inhibits <b>PI3K</b> with IC _{so} of 13 $\mu$ M.	
Clinical Data: No Development Reported Size: 1 mg, 5 mg		Clinical Data:       No Development Reported         Size:       5 mg, 10 mg, 50 mg	
NU9056	<b>Cat. No.</b> : HY-110127	NUN82647 (QBS)	<b>Cat. No.:</b> HY-115683
NU9056 is a potent and selective <b>Tip60 (KAT5)</b> histone acetyltransferase inhibitor with an of 2 $\mu$ M. NU9056 shows >16-fold selectivity for <b>Tip60</b> over PCAF, p300 and GCN5. NU9056 induces <b>apoptosis</b> of prostate cancer cells.	N.S.S.S.N	NUN82647 inhibits cell cycle at G2 phase and induces apoptosis.	NH2 O S NH O
Purity:98.81%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg		Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	
Nutlin-3a		NVP 231	
(Rebemadlin)	<b>Cat. No.</b> : HY-10029		Cat. No.: HY-13945
Nutlin-3a (Rebemadlin), an active enantiomer of Nutlin-3, is a potent <b>murine double minute</b> (MDM2) inhibitor ( $IC_{so}=90$ nM). Nutlin-3a inhibits MDM2-p53 interactions and stabilizes the p53 protein, and induces cell <b>autophagy</b> and <b>apoptosis</b> . Purity: 98.07% Clinical Data: No Development Reported		NVP 231 is a potent, specific, and reversibleceramide kinase (CerK) inhibitor(ICs0=12 nM)that competitively inhibits binding of ceramide toCerK. NVP 231 induces cell apoptosis by increasingDNA fragmentation and caspase-3 and caspase-9cleavage.Purity:98.91%Clinical Data:No Development Reported	Processi
Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg		Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 500 mg	1
NVP-2	<b>Cat. No.</b> : HY-12214A	NVP-ADW742 (ADW742; GSK 552602A; ADW)	<b>Cat. No.:</b> HY-10252
NVP-2 is a potent and selective ATP-competitive cyclin dependent kinase 9 (CDK9) probe, inhibits CDK9/CycT activity with an IC _{s0} of 0.514 nM. NVP-2 displays inhibitory effcts on CDK1/CycB, CDK2/CycA and CDK16/CycY kinases with IC _{s0} values of 0.584 $\mu$ M, 0.706 $\mu$ M, and 0.605 $\mu$ M, respectively.Purity:99.12%Clinical Data:No Development Reported	J. L. T. C.	NVP-ADW742 (ADW742) is an orally active, selectiveIGF-1R tyrosine kinase inhibitor with an $IC_{s0}$ of $0.17 \ \mu$ M. NVP-ADW742 inhibits insulin receptor (InsR)with an $IC_{s0}$ of 2.8 $\mu$ M. NVP-ADW742 inducespleiotropic antiproliferative/proapoptotic biologicsequelae in tumor cells.Purity:99.30%Clinical Data:No Development Reported	N N NH2 COCO
Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	00 mg	Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	

NVP-HSP990		NVP-TAE 226	
(HSP-990)	Cat. No.: HY-15190	(TAE226)	Cat. No.: HY-13203
NVP-HSP990 is a potent and selective Hsp90 inhibitor, with IC _{so} values of 0.6, 0.8, and 8.5 nM for Hsp90α, Hsp90β, and Grp94, respectively. Purity: 99.77% Clinical Data: Phase 1		NVP-TAE 226 (TAE226) is a potent and         ATP-competitive dual FAK and IGF-1R inhibitor with         IC ₅₀ S of 5.5 nM and 140 nM, respectively. NVP-TAE         226 (TAE226) also effectively inhibits Pyk2 and         insulin receptor (InsR) with IC ₅₀ S of 3.5 nM and 44         nM, respectively.         Purity:       99.92%         Clinical Data:       No Development Reported	
Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	.00 mg	Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
NVP-TAE 684		NVP-TNKS656	
(TAE 684)	Cat. No.: HY-10192	(TNKS656)	Cat. No.: HY-13990
NVP-TAE 684 (TAE 684) is a highly potent and selective <b>ALK</b> inhibitor, which blocks the growth of ALCL-derived and ALK-dependent cell lines with <b>IC</b> ₅₀ values between 2 and 10 nM.		NVP-TNKS656 is a highly potent, selective, and orally active TNKS2 inhibitor with $IC_{50}$ of 6 nM, and is > 300 fold selectivity against PARP1 and PARP2.	dir forde
Purity: 99.42%	N	Purity: 99.52%	
Clinical Data: No Development Reported Size: 10 mM × 1 mL 5 mg. 10 mg. 50 mg. 100 mg.		Clinical Data: No Development Reported Size: 10 mM × 1 mL 2 ma, 5 ma, 10 ma, 25 ma, 50	ma. 100 ma
			3, 3
NVS-CECR2-1		NVX-207	
	Cat. No.: HY-110374		Cat. No.: HY-101597
NVS-CECR2-1, a non-BET family <b>Bromodomain</b> ( <b>BRD</b> ) inhibitor, is a potent and selective cat eye syndrome chromosome region, candidate 2 ( <b>CECR2</b> ) inhibitor. NVS-CECR2-1 binds to CECR2 BRD with high affinity ( $IC_{50}$ =47 nM; $K_{D}$ =80 nM).Purity: $\geq 99.0\%$ Clinical Data: Size:Size:5 mg	~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~	NVX-207, a Betulinic acid-derived anti-cancer compound, shows anti-tumor activity (mean $IC_{50}=3.5$ $\mu$ M) against various human and canine cell lines. NVX-207-induced <b>apoptosis</b> is associated with activation of the intrinsic apoptotic pathway via cleavage of caspases -9, -3, -7 and of PARP.Purity: $\geq$ 98.0%Clinical Data: Size:So Development Reported Size:Size:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
Nystatin	Cat No. UV 17400	O6-Benzylguanine	
Nystatin is an orally active polyene <b>antifungal</b> <b>antibiotic</b> effective against yeast and mycoplasma. Nystatin increases the permeability of plasma membranes to small monovalent ions, including chloridion.		O6-Benzylguanine, a guanine analog, is the DNA repair enzyme O6-alkylguanine-DNA alkyltransferase (MGMT/AGT) inhibitor.	
Purity: 98.29%		Purity: 99.63%	
Clinical Data: Launched Size: 200 mg, 500 mg		Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 25 mg, 50 mg, 100 mg	
			·
OBAA	<b>Cat. No</b> .: HY-101015A	Obacunone	<b>Cat. No.:</b> HY-N0428
OBAA is a potent <b>phospholipase A2 (PLA2)</b> inhibitor with an IC ₅₀ of 70 nM. OBAA blocks Melittin-induced Ca ²⁺ influx in Trypanosoma brucei with an IC ₅₀ of 0.4 $\mu$ M.		Obacunone, isolated from seeds of Marsh White grapefruit, exhibits anti-tumor activity by the induction of apoptosis.	
Purity:     >98%       Clinical Data:     No Development Reported       Size:     5 mg, 10 mg		Purity:99.75%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 50 mg	°7∖

Odoroside A	Cat No HV-N7496	ODQ	<b>Cat No</b> : HY-101255
Odoroside A is an active ingredient extracted from the leaves of Nerium oleander Linn. Odoroside A has anti-cancer activity. Odoroside A could induce apoptosis and cell cycle arrest through ROS/p53 signaling pathway, leading to the tumor cell death. Purity: 98.75% Clinical Data: No Development Reported		ODQ is a potent and selective <b>soluble guanylyl</b> <b>cyclase</b> (sGC, nitric oxide-activated enzyme) inhibitor. ODQ enhances the pro-apoptotic effects of Cisplatin in human mesothelioma cells. <b>Purity:</b> 99.52% <b>Clinical Data:</b> No Development Reported	
Size: 1 mg, 5 mg, 10 mg		Size: 10 mM × 1 mL, 10 mg, 25 mg, 50 mg	
Oenothein B	<b>Cat. No.:</b> HY-N7765	Okadaic acid	<b>Cat. No.:</b> HY-N6785
Oenothein B is a dimeric macrocyclic ellagitannin and has widely pharmacological activities, including antioxidant, anti-inflammatory, antifungal, anti-HCV, and antitumor properties. Oenothein B is a potent and specific inhibitor of <b>poly(ADP-ribose) glycohydrolase</b> . <b>Purity:</b> >98% Clinical Data: No Development Reported	and the second	Okadaic acid, a marine toxin, is an inhibitor of protein phosphatases (PP). Purity: ≥98.0% Clinical Data: No Development Reported	Alt Signer for the
Size: 5 mg, 10 mg, 25 mg		Size: 25 μg (124.2 μM * 250 μL in Ethanol)	
Oleic acid (9-cis-Octadecenoic acid; 9Z-Octadecenoic acid)	<b>Cat. No.</b> : HY-N1446	Oleic acid-13C (9-cis-Octadecenoic acid-13C; 9Z-Octadecenoic acid-13C)	<b>Cat. No.:</b> HY-N1446S
Oleic acid (9-cis-Octadecenoic acid) is an abundant monounsaturated fatty acid. Oleic acid is a Na*/K* ATPase activator.	la.	Oleic acid-13C (9-cis-Octadecenoic acid-13C) is the 13C labeled Oleic acid. Oleic acid (9-cis-Octadecenoic acid) is an abundant monounsaturated fatty acid. Oleic acid is a Na*/K* ATPase activator.	
Purity:≥98.0%Clinical Data:LaunchedSize:10 mM × 1 mL, 1 mg, 5 mg, 10 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
Oleic acid-13C-1	<b>Cat. No.</b> : HY-N1446S4	Oleic acid-13C18 (9-cis-Octadecenoic acid-13C18; 9Z-Octadecenoic acid-13C18)	<b>Cat. No.:</b> HY-N1446S2
Oleic acid-13C-1 is the 13C labeled Oleic acid. Oleic acid (9-cis-Octadecenoic acid) is an abundant monounsaturated fatty acid. Oleic acid is a Na ⁺ /K ⁺ ATPase activator.	н.т.,	Oleic acid-13C18 (9-cis-Octadecenoic acid-13C18) is the 13C labeled Oleic acid. Oleic acid (9-cis-Octadecenoic acid) is an abundant monounsaturated fatty acid. Oleic acid is a Na*/K* ATPase activator.	and and a stand and a stand
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
Oleic acid-d17 (9-cis-Octadecenoic acid-d17; 9Z-Octadecenoic acid-d17)	<b>Cat. No.:</b> HY-N1446S3	Oleic acid-d2 (9-cis-Octadecenoic acid-d2; 9Z-Octadecenoic acid-d2)	<b>Cat. No.:</b> HY-N1446S1
Oleic acid-d17 (9-cis-Octadecenoic acid-d17) is the deuterium labeled Oleic acid. Oleic acid (9-cis-Octadecenoic acid) is an abundant monounsaturated fatty acid. Oleic acid is a <b>Na*/K* ATPase</b> activator.	ру ородар р россобран р	Oleic acid-d2 (9-cis-Octadecenoic acid-d2) is the deuterium labeled Oleic acid. Oleic acid (9-cis-Octadecenoic acid) is an abundant monounsaturated fatty acid. Oleic acid is a Na+/K+ ATPase activator.	° → → → → → → → → → → → → →
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	- U






р38 МАРК-IN-3		p53 Activator 2	
	Cat. No.: HY-144697		Cat. No.: HY-146095
p38 MAPK-IN-3 (Compound 2c) is a <b>p38α MAPK</b> inhibitor. p38 MAPK-IN-3 has antitumor activities and induces <b>apoptosis</b> and <b>ROS</b> .	Stora, Star	p53 Activator 2 (compound 10ah) intercalats into DNA and results in significant DNA double-strand break.p53 Activator 2 increases the expression of <b>p53</b> , p-p53, CDK4, p21 to cause cell cycle arrest at G2/M phase.	o ^R Class Harrison
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
PAC-1		Paclitaxel	
(Procaspase activating compound 1)	Cat. No.: HY-13523		Cat. No.: HY-B0015
PAC-1 is a <b>procaspase-3</b> activator that induces apoptosis in cancer cells with an $EC_{s0}$ of 2.08 $\mu$ M.	orou _t o.	Paclitaxel is a naturally occurring antineoplastic agent and stabilizes <b>tubulin</b> polymerization. Paclitaxel can cause both mitotic arrest and <b>apoptotic</b> cell death. Paclitaxel also induces <b>autophagy</b> .	
Purity:         99.93%           Clinical Data:         Phase 2           Size:         10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 500 mg//line	g	Purity:         99.97%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 50 mg, 100 mg, 500 mg	Ó
Paclitaxel-d5 (benzoyloxy)		Paederosidic acid	
	Cat. No.: HY-B0015S1		Cat. No.: HY-N6998
Paclitaxel-d5 benzoyloxy is the deuterium labeled Paclitaxel. Paclitaxel is a naturally occurring antineoplastic agent and stabilizes <b>tubulin</b> polymerization. Paclitaxel can cause both mitotic arrest and <b>apoptotic</b> cell death. Paclitaxel also induces <b>autophagy</b> . Purity: >98% Clinical Data: No Development Reported Size: 1 are 5 are		Paederosidic acid is isolated from P. scandens with anticancer and antiinflammation activities. Paederosidic acid inhibits lung caner cells via inducing mitochondria-mediated <b>apoptosis</b> . <b>Purity:</b> 99.90% <b>Clinical Data:</b> No Development Reported	
Size. I mg, 5 mg		5 mg, 10 mg, 25 mg	
D (0)			
Paeonifiorigenone		PAK4-IN-2	
	Cat. No.: HY-N3119		Cat. No.: HY-143490
Paeoniflorigenone, isolated as an active ingredient from the root of moutan cortex, induces <b>apoptosis</b> selectively in the cancer cell lines and exhibits antiproliferative effect.	o C	PAK4-IN-2 is a highly potent <b>PAK4</b> inhibitor with $IC_{s0}$ value of 2.7 nM. PAK4-IN-2 can arrest MV4-11 cells at G0/G1 phase and induce cell <b>apoptosis</b> . PAK4-IN-2 can be used for researching cancer.	
	(13)		H ₂ N
Purity:     ≥98.0%       Clinical Data:     No Development Reported       Size:     1 mg	0~~_0	Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	2
Palomid 529	Cat No: HV-14581	pan-HER-IN-1	<b>Cat No</b> : HY-144676
	Suc. 10. 111 17501		
Palomid 529 is a potent inhibitor of mTORC1 and mTORC2 complexes.	~ origina	pan-HER-IN-1 (Compound C5) is an irreversible, orally active pan-HER inhibitor with IC ₅₀ values of 0.38, 1.6, 2.2 and 3.5 nM against EGFR, HER4, EGFR ^{T790MI358R} and HER2, respectively. pan-HER-IN-1 induces <b>apoptosis</b> and shows antitumor activities.	
Purity: 99.47%		Purity: >98%	
Clinical Data: Phase 1 Size: 10 mM x 1 ml 5 mg 10 mg 50 mg 100 mg		Clinical Data: No Development Reported	
2.20. 10 mm ·· 1 mz, 3 mg, 10 mg, 30 mg, 100 mg		1 mg, 3 mg	

pan-HER-IN-2		Pan-Trk-IN-3	
	Cat. No.: HY-144677		Cat. No.: HY-144069
pan-HER-IN-2 (Compound C6) is a reversible, orally active pan-HER inhibitor with IC _{so} values of         0.72, 2.0, 8.2 and 75.1 nM against EGFR, HER4,         EGFR ^{T790ML858R} and HER2, respectively.         pan-HER-IN-2 induces apoptosis and shows antitumor activities.         Purity:       >98%         Clinical Data:       No Development Reported         Size:       1 mg, 5 mg		Pan-Trk-IN-3 (Compound 11g) is a potent inhibitor of pan-Trk and their drug-resistant mutants with IC _{\$0} values of 2, 3, 2, 21, 26, 5, 7 and 6 nM against TrkA, TrkB, TrkC, TrkA ^{GS95R} , TrkA ^{G667C} , TrkA ^{G6575} , TrkA ^{G6575} , TrkA ^{G6575} , TrkA ^{G6578} , respectively.         Purity:       >98%         Clinical Data:       No Development Reported         Size:       1 mg, 5 mg	
Panepoxydone	<b>Cat. No.</b> : HY-N10266	Panobinostat (LBH589; NVP-LBH589)	<b>Cat. No.:</b> HY-10224
Panepoxydone is an inhibitor of <b>NF-kB</b> activation. Panepoxydone interferes with the NF-kB mediated signal transduction by inhibiting the phosphorylation of IkB. Panepoxydone exhibits antitumor, anti-inflammatory, antimalarial and anti-parasitic activity. <b>Purity:</b> >98% <b>Clinical Data:</b> No Development Reported <b>Size:</b> 1 mg, 5 mg	OH H OH OH	Panobinostat (LBH589; NVP-LBH589) is a potent and orally active non-selective HDAC inhibitor, and has antineoplastic activities.         Purity:       99.20%         Clinical Data:       Launched         Size:       10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg	, 500 mg
Panobinostat-d4 (LBH589-d4; NVP-LBH589-d4)	<b>Cat. No.</b> : HY-10224S	Panobinostat-d4 hydrochloride (LBH589-d4 hydrochloride; NVP-LBH589-d4 hydrochloride)	<b>Cat. No.</b> : HY-10224S1
Panobinostat-d4 (LBH589-d4) is the deuterium labeled Panobinostat. Panobinostat (LBH589; NVP-LBH589) is a potent and orally active non-selective HDAC inhibitor, and has antineoplastic activities. Purity: >98% Clinical Data: No Development Reported	°fff ⊳fff HN of OH	Panobinostat-d4 (hydrochloride) is deuterium labeled Panobinostat. Panobinostat (LBH589; NVP-LBH589) is a potent and orally active non-selective HDAC inhibitor, and has antineoplastic activities. Purity: >98% Clinical Data: No Development Reported	CHT Ha
Size: 1 mg, 5 mg		Size: 1 mg, 5 mg	
Pantoprazole (BY1023; SKF96022)	<b>Cat. No.</b> : HY-17507	Pantoprazole sodium (BY1023 sodium; SKF96022 sodium)	<b>Cat. No.:</b> HY-17507A
Pantoprazole (BY10232) is an orally active and potent <b>proton pump</b> inhibitor (PPI). Pantoprazole, a substituted benzimidazole, is a potent $H^*/K^*$ - <b>ATPase</b> inhibitor with an IC ₅₀ of 6.8 $\mu$ M.	F→ ^Q →()↓↓→ ^Q → ^Q → ^Q →	Pantoprazole sodium (BY10232 sodium) is an orally active and potent <b>proton pump</b> inhibitor (PPI). Pantoprazole sodium, a substituted benzimidazole, is a potent H*/K*-ATPase inhibitor with an IC _{so} of 6.8 $\mu$ M.	Na [*] -0 0-
Purity:>98%Clinical Data:LaunchedSize:1 mg, 5 mg		Purity:         99.89%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 100 mg, 500 mg	
Pantoprazole sodium hydrate (BY1023 sodium hydrate; SKF96022 sodium hydrate)	<b>Cat. No.:</b> HY-17507B	Pantoprazole-d3 (BY1023-d3; SKF96022-d3)	<b>Cat. No.</b> : HY-17507S1
Pantoprazole sodium hydrate (BY10232 sodium hydrate) is an orally active and potent <b>proton</b> <b>pump</b> inhibitor (PPI). Pantoprazole sodium hydrate, a substituted benzimidazole, is a potent H*/K*-ATPase inhibitor with an IC _{so} of 6.8 μM.		Pantoprazole-d3 is deuterium labeled Pantoprazole. Pantoprazole (BY10232) is an orally active and potent proton pump inhibitor (PPI). Pantoprazole, a substituted benzimidazole, is a potent H+/K+-ATPase inhibitor with an IC50 of 6.8 µM.	
Purity:         99.94%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 100 mg, 500 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	

Pantoprazole-d6		PAO-Nan	
(BY1023-d6; SKF96022-d6)	Cat. No.: HY-17507S		Cat. No.: HY-D1267
Pantoprazole-d6 is deuterium labeled Pantoprazole.         Pantoprazole (BY10232) is an orally active and         potent proton pump inhibitor (PPI). Pantoprazole,         a substituted benzimidazole, is a potent         H+/K+-ATPase inhibitor with an IC50 of 6.8 μM.         Purity:       >98%         Clinical Data:       No Development Reported         Size:       1 mg, 5 mg		PAO-Nap is the modified PAO attached a naphthalimide fluorophore using aminocaproic acid as a linker. PAO induces oxidative stress-mediated apoptosis in HL-60 cells by selectively targeting thioredoxin reductase. br/>.Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg	CH CH AMOH
Paris saponin VII		PARP/PI3K-IN-1	
(Chonglou Saponin VII)	Cat. No.: HY-N3584		Cat. No.: HY-133124
Paris saponin VII (Chonglou Saponin VII) is a steroidal saponin isolated from the roots and rhizomes of Trillium tschonoskii Maxim. Paris saponin VII-induced apoptosis in K562/ADR cells is associated with Akt/MAPK and the inhibition of P-gp. Purity: 99.13%		PARP/PI3K-IN-1 (compound 15) is a potent PARP/PI3K inhibitor with pIC_{s0} values of 8.22, 8.44, 8.25, 6.54, 8.13, 6.08 for PARP-1, PARP-2, PI3Kα, PI3Kβ, PI3Kδ, and PI3Kγ, respectively. PARP/PI3K-IN-1 is a highly effective anticancer compound targeted against a wide range of oncologic diseases.Purity:>98%	gratoža"
Clinical Data: No Development Reported Size: 5 mg, 10 mg		Clinical Data: No Development Reported Size: 1 mg, 5 mg	
PARP1/2/TNKS1/2-IN-1		PARP1/BRD4-IN-1	
	Cat. No.: HY-146336		Cat. No.: HY-144338
PARP1/2/TNKS1/2-IN-1 (Compound I-9) is a dual PARP-1, PARP-2, TNKS1 and TNKS2 inhibitor with IC ₅₀ values of 0.25 nM, 1.2 nM, 13.5 nM and 4.15 nM against PARP-1, PARP-2, TNKS1 and TNKS2, respectively.		PARP1/BRD4-IN-1 is a potent and high selective PARP1/BRD4 inhibitor (IC _{sp} s of 49 and 202 nM in PARP1 and BRD4, respectively). PARP1/BRD4-IN-1 represses the expression and activity of PARP1 and BRD4 to synergistically inhibit the malignant growth of pancreatic cancer cells.	
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg		Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	
PARP10/15-IN-2		PARP10/15-IN-3	
	Cat. No.: HY-146501	1744 20,20 14 0	Cat. No.: HY-146502
PARP10/15-IN-2 (Compound 8h) is a potent PARP10 and PARP15 dual inhibitor with IC ₅₀ values of 0.15 μM and 0.37 μM against PARP10 and PARP15, respectively. PARP10/15-IN-2 is able to enter cells and rescue cells from <b>apoptosis</b> . Purity: >98% Clinical Data: No Development Reported	C F	PARP10/15-IN-3 (Compound 8a) is a potent PARP10 and PARP15 dual inhibitor with IC ₅₀ values of 0.14 μM and 0.40 μM against PARP10 and PARP15, respectively. PARP10/15-IN-3 is able to enter cells and rescue cells from apoptosis.         Purity:       >98%         Clinical Data:       No Development Reported	NH C
Size: 1 mg, 5 mg		Size: 1 mg, 5 mg	
PARP14 inhibitor H10	<b>Cat. No.:</b> HY-117889	Parthenolide ((-)-Parthenolide)	<b>Cat. No.:</b> HY-N0141
PARP14 inhibitor H10, compound H 10, is a selective inhibitor against PARP14 ( $IC_{s_0}$ =490 nM), over other PARPs ( $\approx$ 24 fold over PARP1). PARP14 inhibitor H10 induces caspase-3/7-mediated cell apoptosis.	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Parthenolide is a sesquiterpene lactone found in the medicinal herb Feverfew. Parthenolide exhibits anti-inflammatory activity by inhibiting NF-κB activation; also inhibits HDAC1 protein without affecting other class I/II HDACs.	
Purity:     98.16%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg, 10 mg		Purity:         99.13%           Clinical Data:         Phase 2           Size:         10 mM × 1 mL, 50 mg, 100 mg, 200 mg	ŏ

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Patulin (Terinin)	Cat No : HV-N6779	PB28	Cat No : HV_1085114
Patulin (Terinin) is a mycotoxin produced by fungi including the Aspergillus, Penicillium, and Byssochlamys species, is suspected to be clastogenic, mutagenic, teratogenic and cytotoxic.		PB28 is a cyclohexylpiperazine derivative and a high affinity and selective <b>sigma 2 (o2)</b> <b>receptor</b> agonist with a $K_i$ of 0.68 nM. PB28 is also a $\sigma$ 1 antagonist with a $K_i$ of 0.38 nM. PB28 is less affinity for other receptors.	
Purity:99.47%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 25 mg		Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	, o
PB28 dihydrochloride	<b>Cat. No.:</b> HY-108511	PBOX 6	<b>Cat. No.:</b> HY-U00446
PB28 dihydrochloride, a cyclohexylpiperazine derivative, is a high affinity and selective <b>sigma 2 ($\sigma$2) receptor</b> agonist with a <b>K</b> _i of 0.68 nM. PB28 dihydrochloride is also a $\sigma$ 1 antagonist with a <b>K</b> _i of 0.38 nM.	H-CI	PBOX 6 is a pyrrolo-1,5-benzoxazepine (PBOX) compound, acts as a <b>microtubule</b> -depolymerizing agent and an apoptotic agent.	
Purity:99.53%Clinical Data:No Development ReportedSize:5 mg, 10 mg	_о н-а	Purity:99.68%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg	
PCC0208017	<b>Cat. No.</b> : HY-139604	PCI-34051	<b>Cat. No.:</b> HY-15224
PCC0208017 is a microtubule affinity regulating kinases (MARK3/MARK4) inhibitor with $IC_{so}s$ of 1.8 and 2.01 nM, respectively. PCC0208017 has much lower inhibitory activity against MARK1 and MARK2, with $IC_{so}s$ of 31.4 and 33.7 nM, respectively.	HALL HALL	PCI-34051 is a potent and selective HDAC8 inhibitor with $IC_{50}$ of 10 nM, with >200-fold selectivity over the other HDAC isoforms.	OH CTN COO
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg		Purity:         99.64%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
PD-1/PD-L1-IN-10	<b>Cat. No.:</b> HY-132202	PD0166285	<b>Cat. No.:</b> HY-13925
PD-1/PD-L1-IN-10 (compound B2) is an orally active <b>PD-1/PD-L1</b> inhibitor ( <b>IC</b> _{s0} of 2.7 nM) with potent anticancer efficacy.	C C C C C C C C C C C C C C C C C C C	PD0166285, a substrate of P-gp, is a WEE1 inhibitor and a weak Myt1 inhibitor with $IC_{s_0}$ values of 24 and 72 nM, respectively. PD0166285 exhibits an $IC_{s_0}$ of 3.433 $\mu$ M for Chk1.	HN N N N O
Purity:99.29%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg		Purity:         99.46%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	0 mg
PD0166285 dihydrochloride	<b>Cat. No</b> .: HY-13925A	PD168393	<b>Cat. No.</b> : HY-13896
PD0166285 dihydrochloride, a substrate of P-gp, is a WEE1 inhibitor and a weak Myt1 inhibitor with IC ₅₀ values of 24 and 72 nM, respectively. PD0166285 dihydrochloride exhibits an IC ₅₀ of 3.433 $\mu$ M for Chk1.	HN N N O O	PD168393 is a potent, selective and cell-permeable inhibitor of EGFR tyrosine kinase and ErbB2. PD168393 irreversiblely inactivates EGF receptor ( $IC_{so}$ =0.7 nM) and is inactive against insulin receptor, PDGFR, FGFR and PKC.	
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	́,×́	Purity:         98.60%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 25 mg, 50 mg	Br

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PD1/30/4	<b>Cat No</b> : HY-10321	PD173955	<b>Cat No</b> : HY-10395
PD173074 is a potent FGFR1 inhibitor with an $IC_{50}$ of 25 nM and also inhibits VEGFR2 with an $IC_{50}$ of 100-200 nM, showing 1000-fold selectivity for FGFR1 over PDGFR and c-Src.		PD173955 is src family-selective tyrosine kinase inhibitor with IC50 of ~22 nM for Src, Yes and Abl kinase; less potent for FGFR $\alpha$ and no activity on InsR and PKC.	
Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 m	g	Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 5 mg, 10 mg	
PD180970	<b>Cat. No.:</b> HY-103274	PD184161	Cat. No.: HY-10174
PD180970 is a highly potent and ATP-competitive $p210^{Ber-Abl}$ kinase inhibitor, with an $IC_{s0}$ of 5 nM for inhibiting the autophosphorylation of $p210^{Ber-Abl}$ . PD180970 also inhibits Src and KIT kinase with $IC_{s0}$ s of 0.8 nM and 50 nM, respectively. Purity: >98%		PD184161 is an orally active <b>MEK</b> inhibitor. PD184161 inhibits MEK activity (IC ₅₀ =10-100 nM) in a time- and concentration-dependent manner. PD184161 inhibits cell proliferation and induces <b>apoptosis</b> . PD184161 produces depressive-like behavior. <b>Purity:</b> 99.38%	
Clinical Data:         No Development Reported           Size:         5 mg, 10 mg		Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
PDGFR-IN-1	Cat. No : HV.144653	PDK4-IN-1	Cat No : HV-135054
PDGFR-IN-1 (compound 7m) is a potent and orally active PDGFR (platelet-derived growth factor receptor) inhibitor, with IC _{so} values of 2.4 and 0.9 nM for PDGFRα and PDGFRβ, respectively.         Purity:       >98%         Clinical Data:       No Development Reported		PDK4-IN-1 is an anthraquinone derivative and a potent and orally active <b>pyruvate dehydrogenase</b> <b>kinase 4 (PDK4)</b> inhibitor with an IC ₅₀ value of 84 nM. PDK4-IN-1 potently represses cellular transformation and cellular proliferation and induces <b>apoptosis</b> . <b>Purity:</b> >98% <b>Clinical Data:</b> No Development Reported	
Size: 1 mg, 5 mg		Size: 1 mg, 5 mg	
PDK4-IN-1 hydrochloride	<b>Cat. No.:</b> HY-135954A	PDPOB	Cat. No.: HY-145243
PDK4-IN-1 hydrochloride is an anthraquinone derivative and a potent and orally active <b>pyruvate dehydrogenase kinase 4 (PDK4)</b> inhibitor with an IC ₅₀ value of 84 nM.		PDPOB is a phenyl carboxylic acid derivative. PDPOB displays protective roles against OGD/R-evoked multiaspect neuronal deterioration in SH-SY5Y cells, as evidenced by alleviated mitochondrial dysfunction, oxidative stress, and <b>apoptosis</b> .	HOLICIA
Purity:99.48%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	
PEAQX tetrasodium hydrate (NVP-AAM077 tetrasodium hydrate)	<b>Cat. No.:</b> HY-12294A	Pectolinarin	<b>Cat. No.:</b> HY-N0314
PEAQX (NVP-AAM077) tetrasodium hydrate is a potent, selective and orally active NMDA antagonist, with $IC_{50}$ values of 270 nM and 29600 nM for hNMDAR 1A and hNMDAR 2A, respectively.	Br H O'CNa H O'CNa	Pectolinarin possesses anti-inflammatory activity. Pectolinarin inhibits secretion of IL-6 and IL-8, as well as the production of PGE2 and NO. Pectolinarin suppresses cell proliferation and inflammatory response and induces <b>apoptosis</b> via inactivation of the <b>PI3K/Akt</b> pathway.	HQ CH OF OF
Purity:97.05%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg		Purity:99.89%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 20 mg	ности

Pelcitoclax		Pemetrexed disodium	
(APG-1252)	Cat. No.: HY-109185	(LY231514 disodium)	Cat. No.: HY-10820A
Pelcitoclax (APG-1252) is a potent Bcl-2/Bcl-xl inhibitor with antineoplastic and pro-apoptotic effects.         Purity:       95.53%         Clinical Data:       Phase 2         Size:       1 mg, 5 mg, 10 mg, 25 mg, 50 mg	2000 0000 miles	Pemetrexed disodium (LY231514 disodium) is an antifolate, the K _i s of the pentaglutamate of Pemetrexed disodium are 1.3, 7.2, and 65 nM for inhibits thymidylate synthase (TS), dihydrofolate reductase (DHFR), and glycinamide ribonucleotide formyltransferase (GARFT), respectively.         Purity:       99.23%         Clinical Data:       Launched         Size:       10 mM × 1 mL, 50 mg, 100 mg, 200 mg	HAN THE PART OF TH
Pemetrexed disodium hemipenta hydrate (LY231514 disodium hemipenta hydrate)	<b>Cat. No</b> .: HY-13781	Pemetrexed-d5 disodium (LY231514-d5 disodium)	Cat. No.: HY-10820AS
Pemetrexed disodium hemipenta hydrate is a novel antifolate, the K _i values of the pentaglutamate of LY231514 are 1.3, 7.2, and 65 nM for inhibits thymidylate synthase (TS), dihydrofolate reductase (DHFR), and glycinamide ribonucleotide formyltransferase (GARFT), respectively.		Pemetrexed-d5 (LY231514-d5) disodium is the deuterium labeled Pemetrexed disodium.	
Purity:         99.89%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 100 mg, 500 mg		Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	
	]		
Penicillic acid		Pentagamavunon-1	C . N. UV 100477
	Cat. NO.: HY-N6777	(FGV-I)	Cat. NO.: HY-1364/7
Penicillic acid is a polyketide mycotoxin produced by several species of Aspergillus and Penicillium. Penicillic acid exhibits cytotoxicity in rat alveolar macrophages (AM) in vitro.	ОН ОН	Pentagamavunon-1 (PGV-1), a Curcumin analog with oral activity, targets on several molecular mechanisms to induce <b>apoptosis</b> including inhibition of angiogenic factors cyclooxygenase-2 ( <b>COX-2</b> ) and vascular endothelial growth factor ( <b>VEGF</b> ). PGV-1 inhibits <b>NF-κB</b> activation.	HO C C C C C C C C C C C C C C C C C C C
Purity:99.83%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg		Purity:99.80%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg, 50 mg, 100 mg	
Perifosine		Perillyl alcohol	
(KRX-0401; NSC 639966; D21266)	Cat. No.: HY-50909		Cat. No.: HY-N7000
Perifosine is an oral <b>Akt</b> inhibitor which inhibits proliferation of different tumor cell lines with $IC_{50}$ s of 0.6-8.9 µM.	~~~~~~ _* \$ _* ° [*]	Perillyl alcohol, a monoterpene, is active in inducing <b>apoptosis</b> in tumor cells without affecting normal cells.	Сон
Purity: ≥98.0%		<b>Purity:</b> ≥95.0%	
Clinical Data:         Phase 3           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Clinical Data: Phase 2 Size: 100 mg	
Perindopril erbumine (Perindopril tert-butylamine salt; S-9490 erbumine)	Cat. No.: HY-B0130A	Periplocin	Cat. No.: HY-N1381
Perindopril erbumine (Perindopril tert-butylamine salt) is a potent ACE inhibitor of which is used to treat high blood pressure, heart failure or stable coronary artery disease. Target: ACE Perindopril is a long-acting ACE inhibitor.	H H ON HNH	Periplocin is a cardiotonic steroid isolated from Periploca forrestii. Periplocin promotes tumor cell <b>apoptosis</b> and inhibits tumor growth. Periplocin has the potential to facilitate wound healing through the activation of Src/ERK and PI3K/Akt pathways mediated by Na/K-ATPase.	
Purity:         99.98%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 100 mg, 500 mg		Purity:99.79%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 20 mg	

PETCM       Cat. No.: HY-103349       Petromurin C       Cat. No.: HY-10349         PETCM is an activator of caspase-3 and acts as an cytochrome c (cyto c)-dependent manner. PETCM promotes Apaf-1 oligomerization and induces cell apoptosis in HeLa cells.	N10221
PETCM is an activator of <b>caspase-3</b> and acts as an cytochrome c (cyto c)-dependent manner. PETCM promotes Apaf-1 oligomerization and induces cell apoptosis in HeLa cells. N OH	
~	
Purity:     99.36%     Purity:     >98%       Clinical Data:     No Development Reported     Clinical Data:     No Development Reported       Size:     10 mM × 1 mL, 5 mg, 10 mg     Size:     1 mg, 5 mg	Ň.
Pexidartinib     Pexidartinib hydrochloride       (PLX-3397)     Cat. No.: HY-16749       (PLX-3397 hydrochloride)     Cat. No.: HY-16749	16749A
Pexidartinib (PLX-3397) is a potent, orally active, selective, and ATP-competitive colony stimulating factor 1 receptor (CSF1R or M-CSFR) and c-Kit inhibitor, with IC ₅₀ s of 20 and 10 nM, respectively. Pexidartinib hydrochloride (PLX-3397 hydrochloride) is a potent, orally active, selective, and ATP-competitive colony stimulating factor 1 receptor (CSF1R or M-CSFR) and c-Kit inhibitor, with IC ₅₀ s of 20 and 10 nM, respectively.	a Arte
Purity:         99.64%         Purity:         99.89%           Clinical Data:         Launched         Clinical Data:         Launched           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg         Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 200 mg, 500 mg, 1 g	
PF-3758309         PF-3758309 dihydrochloride           (PF-03758309)         Cat. No.: HY-13007         (PF-03758309 dihydrochloride)         Cat. No.: HY-	13007B
PF-3758309 (PF-03758309) is a potent, orally available, and reversible ATP-competitive inhibitor of PAK4 (K _d = 2.7 nM; K _i =18.7 nM).	
Purity:     98.52%     Purity:     >98%       Clinical Data:     No Development Reported     Clinical Data:     No Development Reported       Size:     10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg     Size:     1 mg, 5 mg	1
PF-3/58309 hydrochloride PF-4989216	-13864
PF-3758309 (PF-03758309) hydrochloride is a potent, orally available, and reversible     PF-4989216 is a potent and selective PI3Kα inhibitor with a K ₁ of 0.6 nM.       ATP-competitive inhibitor of PAK4 (K _d = 2.7 nM;     Image: Cut Hout Hout Hout Hout Hout Hout Hout Ho	
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg         Purity:     99.69%       Clinical Data:     No Development Reported       Size:     10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
DE 542 Citrata	
(Sphingosine Kinase 1 Inhibitor II) Cat. No.: HY-15425 (Sphingosine Kinase 1 Inhibitor II Citrate) Cat. No.: HY-	15425A
PF-543 (Sphingosine Kinase 1 Inhibitor II) is a potent, selective, reversible and sphingosine-competitive SPHK1 inhibitor with an IC ₅₀ of 2 nM and a K ₁ of 3.6 nM. PF-543 is >100-fold selectivity for SPHK1 over SPHK2.	CL N HO
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg         Purity:     98.35%       Clinical Data:     No Development Reported       Size:     10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	22.12

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# PI-273

# Cat. No.: HY-103489

PI-273 is a first reversibly and specific phosphatidylinositol 4-kinase (PI4KIIα) inhibitor with an  $I\!C_{_{50}}$  of 0.47  $\mu\text{M}.$  PI-273 can inhibit breast cancer cell proliferation, block the cell cycle and induce cell apoptosis.

Cat. No.: HY-146751

Purity: Clinical Data: No Development Reported Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

# PI3K/Akt/mTOR-IN-2

>98.0%

PI3K/Akt/mTOR-IN-2 is a PI3K/AKT/mTOR pathway inhibitor. PI3K/Akt/mTOR-IN-2 possess anti-cancer effects and selectivity against MDA-MB-231 cells with  $IC_{so}$  value of 2.29  $\mu M.$  PI3K/Akt/mTOR-IN-2 can induce cancer cell cycle arrest and apoptosis

Purity: > 98% Clinical Data: No Development Reported Size: 1 mg, 5 mg

# **ΡΙ3Κδ-ΙΝ-11**

### PI3K $\delta$ -IN-11 is a highly potent and selective PI3K $\delta$ inhibitor with $IC_{50}$ value of 27.5 nM. PI3K $\delta$ -IN-11 dose-dependently blocks the activity of PI3K/Akt pathway. PI3Kδ-IN-11 can be used for researching B or T cell-related malignancies.

>98% Purity: Clinical Data: No Development Reported Size: 1 mg, 5 mg

## Piclidenoson (IB-MECA; CF-101)

Piclidenoson (IB-MECA) is a first-in-class, orally active and selective A3 adenosine receptor (A3AR) agonist. Piclidenoson exhibits antiproliferative effect and induces apoptosis in different cancer cell types like melanoma, leukemia.

99.32% Purity: Clinical Data: Phase 3  $10~\text{mM}\times1$  mL, 5 mg, 10 mg, 25 mg, 50 mg Size

# Picrocrocin

### Picrocrocin, an apocarotenoid found in the flowers of Cochliobolus sativus. Picrocrocin shows anticancer effect. Picrocrocin exhibits growth inhibitory effects against SKMEL-2 human malignant melanoma cells.

Purity: 99.93% Clinical Data: No Development Reported Size: 1 mg, 5 mg



Cat. No.: HY-N4114

### PI3Kγ, PI3Kδ and AKT, respectively). PI3K/AKT-IN-1 has anticancer activity and acts by inhibiting PI3K/AKT axis and inducing caspase 3 dependent

apoptosis. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg

PI3K/AKT-IN-1 is an effective PI3K/AKT dual

PI3Kδ-IN-10 is a highly potent and orally active PI3Kδ inhibitor with IC₅₀ of 2 nM. PI3Kδ-IN-10

induce subsequent apoptosis in hepatocellular

robustly suppresses the downstream AKT pathway to

inhibitor (IC  $_{ro}$  of 6.99, 4.01 and 3.36  $\mu$ M for

# ΡΙ3Κδ-ΙΝ-10

carcinoma models.

**Purity:** 

Size:

PI3K/AKT-IN-1

# **Piceatannol**

# (Astringenin; trans-Piceatannol)

>98% Clinical Data: No Development Reported

1 mg, 5 mg

Piceatannol is a well-known Syk inhibitor and reduces the expression of iNOS induced by TNF. Piceatannol is an effective agent for research of acute lung injury (ALI).



Cat. No.: HY-N9507

98.09% Purity: Clinical Data: No Development Reported Size 10 mM × 1 mL, 10 mg, 25 mg, 50 mg, 100 mg

# **Picrasidine Q**

Picrasidine Q, an alkaloid component extracted from Angelica keiskei species, has the capacity of anti-cell transformation and anti-cancer. Picrasidine Q induces cell apoptosis and G1 phase arrest in human esophageal cancer cell lines, and directly inhibits FGFR2 kinase activity.

Purity: >98% Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg

# Picropodophyllin

# (AXL1717; Picropodophyllin; PPP)

Picropodophyllin (AXL1717) is a selective insulin-like growth factor-1 receptor (IGF-1R) inhibitor with an IC₅₀ of 1 nM.



99.90% Purity: Clinical Data: Phase 3 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg Size:

Cat. No.: HY-13518



Cat. No.: HY-144254

Cat. No.: HY-144806









Picropodophyllotovin-d6		Picroside II	
	Cat. No.: HY-15494S1		Cat. No.: HY-N0408
Picropodophyllotoxin-d6 is deuterium labeled         Picropodophyllin. Picropodophyllin (AXL1717) is a selective insulin-like growth factor-1 receptor (IGF-1R) inhibitor with an IC50 of 1 nM.         Purity:       >98%         Clinical Data:       No Development Reported         Size:       1 mg, 5 mg		Picroside II, an iridoid compound extracted from         Picrorhiza, exhibits anti-inflammatory and         anti-apoptotic activities. picroside II alleviates         the inflammatory response in sepsis and enhances         immune function by inhibiting the activation of         NLRP3 inflammasome and NF-κB pathways.         Purity:       99.77%         Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	но учербан история ан
Pictilisib (GDC-0941)	Cat. No.: HY-50094	Pictilisib dimethanesulfonate (GDC-0941 dimethanesulfonate; GDC-0941 2 MeSO3H salt)	Cat. No.: HY-20180
Pictilisib (GDC-0941) is a potent inhibitor of <b>PI3K$\alpha/\delta$</b> with an <b>IC</b> ₅₀ of 3 nM, with modest selectivity against p110 $\beta$ (11-fold) and p110 $\gamma$ (25-fold).		Pictilisib dimethanesulfonate (GDC-0941 dimethanesulfonate) is a potent inhibitor of PI3K $\alpha/\delta$ with IC _{so} of 3 nM, with modest selectivity against p110 $\beta$ (11-fold) and p110 $\gamma$ (25-fold).	
Purity:         99.80%           Clinical Data:         Phase 2           Size:         10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg	028	Purity:         99.31%           Clinical Data:         Phase 2           Size:         10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg	• •
		DIK 75 hydrochlorida	
PIN-75	<b>Cat. No.:</b> HY-107834	PIK-75 hydrochioride	Cat. No.: HY-13281
PIK-75 is a reversible DNA-PK and p110α-selective inhibitor, which inhibits DNA-PK, p110α and p110γ with IC ₅₆ S of 2, 5.8 and 76 nM, respectively.PIK-75 inhibits p110α > 200-fold more potently than p110β (IC ₅₀ =1.3 $\mu$ M). PIK-75 induces <b>apoptosis</b> .Purity:>98% Clinical Data: Size:1 mg, 5 mg	Br C N N S C N N S C N N S C N S C N S C S	PIK-75 hydrochloride is a reversible DNA-PK and $p110\alpha$ -selective inhibitor, which inhibits DNA-PK, $p110\alpha$ and $p110\gamma$ with $IC_{so}$ of 2, 5.8 and 76 nM,         respectively. PIK-75 hydrochloride inhibits $p110\alpha$ >200-fold more potently than $p110\beta$ (IC _{so} =1.3 $\mu$ M). PIK-75 hydrochloride induces apoptosis.         Purity:       99.72%         Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 10 mg, 50 mg, 100 mg	Br CNN N O N'O
Pilaralisih analogue		PIM-447 dibydrochloride	
(XL147 analogue)	Cat. No.: HY-11105	(LGH447 dihydrochloride)	Cat. No.: HY-19322B
Pilaralisib analogue (XL147 analogue) is a representative and selective PI3Kα inhibitor extracted from patent WO2012006552A1, Compound 147 in Table 1.         Purity:       99.67%         Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 10 mg, 50 mg, 100 mg	U D=S=O NH NH NH N N N	PIM447 dihydrochloride (LGH447 dihydrochloride) is         a potent, orally available, and selective         pan-PIM kinase inhibitor, with K, values of 6,         18, and 9 pM for PIM1, PIM2, and PIM3,         respectively. PIM447 dihydrochloride displays dual         antimyeloma and bone-protective effects.         Purity:       99.27%         Clinical Data:       Phase 1         Size:       10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 50 mg, 100	$ \begin{array}{c} F \\ F \\ F \\ F \\ HCI \\ $
		DIMAAZ	
C-NI-TIME	Cat. No.: HY-143897	(LGH447)	Cat. No.: HY-19322
PIM1-IN-3 (Compound HL8) is a potent inhibitor of <b>PIM1</b> . PIM1-IN-3 shows selective inhibition for the PIM-1 enzyme. PIM1-IN-3 induces apoptosis efficiently in Colo320 cells. PIM1-IN-3 has the potential for the research of cancer diseases.		PIM447 (LGH447) is a potent, orally available, and selective pan- <b>PIM kinase</b> inhibitor, with <b>K</b> ₁ values of 6, 18, and 9 pM for PIM1, PIM2, and PIM3, respectively. PIM447 displays dual antimyeloma and bone-protective effects. PIM447 induces <b>apoptosis</b> .	
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg		Purty:     >98%       Clinical Data:     Phase 1       Size:     1 mg, 5 mg	S7

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Pimpinollin		Pinghanksin	
Finiphienn	Cat. No.: HY-N0438	(3,5,7-Trihydroxyflavanone)	Cat. No.: HY-N3062
Pimpinellin is a constituent of Cyrtomium fortumei (J.). Pimpinellin inhibits the growth of tumor cells via the induction of tumor cell <b>apoptosis</b> .		Pinobanksin has apoptotic induction in a B-cell lymphoma cell line.	но он он
Purity:99.27%Clinical Data:No Development ReportedSize:5 mg, 10 mg	-23/5 <b>N</b>	Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg	81
Pinoresinol ((+)-Pinoresinol)	<b>Cat. No.:</b> HY-N6253	Pinosylvin	<b>Cat. No.:</b> HY-N2387
Pinoresinol is a lignol of plant origin serving for defense in a caterpillar. Pinoresinol drastically sensitizes cancer cells against TNF-related apoptosis-inducing ligand (TRAIL) -induced apoptosis.		Pinosylvin is a pre-infectious stilbenoid toxin isolated from the heartwood of Pinus spp, has <b>anti-bacterial</b> activities. Pinosylvin is a resveratrol analogue, can induce cell <b>apoptosis</b> and <b>autophapy</b> in leukemia cells.	но
Purity:99.69%Clinical Data:No Development ReportedSize:5 mg, 10 mg	J OH	Purity:99.66%Clinical Data:No Development ReportedSize:5 mg, 10 mg	722.044
Pinusolide	<b>Cat. No.:</b> HY-N3055	Piperlongumine (Piplartine)	<b>Cat. No.:</b> HY-N2329
Pinusolide is a known platelet-activating factor (PAF) receptor binding antagonist. Pinusolide not only decreases the proliferation activity of tumor cells but specifically induces <b>apoptosis</b> .		Piperlongumine is a alkaloid, possesses ant-inflammatory, antibacterial, antiangiogenic, antioxidant, antitumor, and antidiabetic activities. Piperlongumine induces ROS, and induces apoptosis in cancer cell lines.	
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	H/ T C	Purity:99.19%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg	
Pitavastatin (NK-104)	<b>Cat. No.:</b> HY-B0144A	Pitavastatin Calcium (NK-104 hemicalcium; Pitavastatin hemicalcium)	<b>Cat. No.:</b> HY-B0144
Pitavastatin (NK-104) is a potent hydroxymethylglutaryl-CoA (HMG-CoA) reductase inhibitor. Pitavastatin inhibits cholesterol synthesis from acetic acid with an $IC_{50}$ of 5.8 nM in HepG2 cells.	PH PH P CLNC CH P	Pitavastatin Calcium (NK-104 hemicalcium) is a potent <b>hydroxymethylglutaryl-CoA (HMG-CoA)</b> reductase inhibitor. Pitavastatin Calcium (NK-104 hemicalcium) inhibits cholesterol synthesis from acetic acid with an $IC_{50}$ of 5.8 nM in HepG2 cells.	
Purity:     >98%       Clinical Data:     Launched       Size:     1 mg, 5 mg		Purity:       99.45%         Clinical Data:       Launched         Size:       10 mM × 1 mL, 10 mg, 50 mg, 100 mg	
Pitavastatin D4 (NK-104 D4)	<b>Cat. No.:</b> HY-B0144AS	Pitavastatin-d4 hemicalcium (NK-104-d4 hemicalcium; Pitavastatin-d4 hemicalcium)	<b>Cat. No.:</b> HY-B0144S
Pitavastatin D4 (NK-104 D4) is deuterium labeled Pitavastatin. Pitavastatin is a potent HMG-CoA reductase inhibitor.		Pitavastatin-d4 (hemicalcium) is deuterium labeled Pitavastatin (Calcium). Pitavastatin Calcium (NK-104 hemicalcium) is a potent hydroxymethylglutaryl-CoA (HMG-CoA) reductase inhibitor.	
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	,	Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	<i></i>

Pitstop 2		Pivanex	
	Cat. No.: HY-115604	(AN-9; Pivalyloxymethyl butyrate)	Cat. No.: HY-120508
Pitstop 2 is a <b>clathrin</b> inhibitor which inhibits <b>clathrin-mediated endocytosis (CME)</b> by associating with the terminal domain of clathrin. Pitstop 2 has the potential for anti-cancer research.		Pivanex (AN-9), a derivative of Butyric acid, is an orally active <b>HDAC</b> inhibitor. Pivanex down-regulates <b>bcr-abl</b> protein and enhances <b>apoptosis</b> . Pivanex has antimetastic and antiangiogenic properties.	
Clinical Data: No Development Reported		Clinical Data: Phase 2	
Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg		Size: 1 mg, 5 mg	
PKCβ inhibitor 1		Pladienolide B	
	Cat. No.: HY-13335		Cat. No.: HY-16399
PKCβ inhibitor 1 is a potent, ATP-competitive, and selective PKCβ inhibitor with IC _{so} s of 21 and 5 nM for human PKCβ1 and PKCβ2, respectively. PKCβ inhibitor 1 exhibits selectivity of more than $60$ -fold in favor of PKCβ2 relative to other PKC isozymes (PKCα, PKCγ, and PKCε).Purity:98.21%Clinical Data Clinical DataDevelopment Paperted	O-H-GN-N	Pladienolide B is a potent cancer cell growth inhibitor that targets the SF3B1 subunit of the spliceosome. Pladienolide B exerts antitumor activities mediated through the inhibition of pre-mRNA splicing. Pladienolide B induces apoptosis. Purity: ≥98.0%	
Size: 500 μg, 1 mg, 5 mg, 10 mg		Size: 100 µg	
		15	
PLK1/BRD4-IN-1		PluriSIn 1	
	Cat. No.: HY-143471	(NSC 14613)	Cat. No.: HY-15700
PLK1/BRD4-IN-1 (9b) is an orally active dual <b>PLK1</b> and <b>BRD4</b> inhibitor with $IC_{s0}$ values of 22 nM and 109 nM against PLK1 and BRD4, respectively.	The state of the s	PluriSIn 1 (NSC 14613) is an inhibitor of stearoyl-coA desaturase (SCD), and is a pluripotent cell-specific inhibitor.	N N N H
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	~	Purity:99.64%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg, 50 mg	
PND-1186		PND-1186 bydrochloride	
(VS-4718: SR-2516)	Cat No : HY-13917	(VS-4718 hydrochloride: SR-2516 hydrochloride)	Cat No : HY-13917A
PND-1186 (VS-4718) is a potent, highly-specific and reversible inhibitor of FAK with an IC _{so} of 1.5 nM. PND-1186 selectively promotes tumor cell <b>apoptosis</b> .		PND-1186 hydrochloride (VS-4718 hydrochloride) is a potent, highly-specific and reversible inhibitor of FAK with an $IC_{so}$ of 1.5 nM. PND-1186 hydrochloride selectively promotes tumor cell <b>apoptosis</b> .	
Purity: 99.80%		Purity: 98.78%	
Clinical Data: Phase 1		Clinical Data: No Development Reported	
Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
PNU-74654	<b>Cat. No.:</b> HY-101130	Podocarpusflavone A	<b>Cat. No.:</b> HY-N2198
PNU-74654 is an inhibitor of $Wnt/\beta\text{-catenin}$ pathway with an $IC_{s0}$ of 129.8 $\mu\text{M}$ in NCI-H295 cell.	Contractor	Podocarpusflavone A is a DNA topoisomerase I inhibitor. Podocarpusflavone A has moderated anti-proliferative activity and induces cell apoptosis in MCF-7. Podocarpusflavone A is developing anti-tumor drugs.	HO LOCATION
Purity:         99.42%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	00 mg, 200 mg	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	он о

Pogostone	Cat. No.: HY-N1416	Polydatin (Piceid)	Cat. No.: HY-N0120A
anti-bacterial and anti-cancer activities.	он он	Polydatin (Piceid), extracted from the roots of Polygonum cuspidatum Sieb, a widely used traditional Chinese remedies, possesses anti-inflammatory activity in several experimental models. Polydatin (Piceid) inhibits G6PD and induces oxidative and ER stresses.	HO TO OUT ON ON ON ON
Purity:99.80%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg		Purity:         98.55%           Clinical Data:         Phase 2           Size:         10 mM × 1 mL, 50 mg, 100 mg, 200 mg, 500 mg	ng
Polygalacin D	<b>Cat. No.:</b> HY-N6064	Polyinosinic-polycytidylic acid sodium (Poly(I:C) sodium)	<b>Cat. No.:</b> HY-135748
Polygalacin D (PGD) is a bioactive compound isolated from Platycodon grandiflorum (Jacq.) with anticancer and anti-proliferative properties.	Stall Barrie	Polyinosinic-polycytidylic acid sodium (Poly(I:C) sodium) is a synthetic analog of double-stranded RNA and an agonist of toll-like receptor 3 (TLR3) and retinoic acid inducible gene I (RIG-I)-like receptors (RIG-I and MDA5).	[zzzie_]zzzieł
Purity:99.30%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:     ≥98.0%       Clinical Data:     No Development Reported       Size:     10 mg, 25 mg	
Polyphyllin G	<b>Cat. No.:</b> HY-N0817	Polyphyllin I	<b>Cat. No.</b> : HY-N0047
Polyphyllin G is isolated from the rhizomes of Paris yunnanensis, with antimicrobial and anticancer activity. Polyphyllin G prevents the growth of both Gram-positive and Gram-negative bacteria with minimum inhibitory concentrations (MICs).	- Storage - Storage	Polyphyllin I is a bioactive constituent extracted from Paris polyphylla, has strong anti-tumor activity. Polyphyllin I is an activator of the JNK signaling pathway and is an inhibitor of PDK1/Akt/mTOR signaling. Polyphyllin I induces autophagy, G2/M phase arrest and apoptosis.	
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg		Purity:99.61%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 20 mg	
Polyphyllin II	<b>Cat. No.:</b> HY-N0048	Polyphyllin VI	<b>Cat. No.:</b> HY-N0816
Polyphyllin II is one of the most significant saponins in Rhizoma Paridis and has toxic effects on kinds of cancer cells. Polyphyllin II induces <b>apoptosis</b> through caspases activation and cell-cycle arrest.		Polyphyllin VI, an active saponin, possess anti-cancer activities. Polyphyllin VI induces G2/M cell cycle arrest and triggers <b>apoptosis</b> .	
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg	Сн	Purity:98.34%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 20 mg	w.s.
Polyporenic acid C	<b>Cat. No</b> .: HY-N2993	Pomalidomide (CC-4047)	<b>Cat. No.:</b> HY-10984
Polyporenic acid C is a lanostane-type triterpenoid isolated from P. cocos. Polyporenic acid C induces cell <b>apoptosis</b> through the death receptor-mediated apoptotic pathway without the involvement of the mitochondria. Polyporenic acid C is promising agent for lung cancer therapy.		Pomalidomide, the third-generation immunomodulatory agent, acts as molecular glue. Pomalidomide interacts with the E3 ligase cereblon and induces degradation of essential Ikaros transcription factors.	
Purity:     ≥99.0%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg		Purity:         99.96%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg	g, 500 mg

Pomalidomide-d3		Pomalidomide-d5	
(CC-4047-d3)	Cat. No.: HY-10984S1	(CC-4047-d5)	Cat. No.: HY-10984S
Pomalidomide-d3 (CC-4047-d3) is the deuterium labeled Pomalidomide. Pomalidomide, the third-generation immunomodulatory agent, acts as molecular glue. Pomalidomide interacts with the E3 ligase cereblon and induces degradation of essential Ikaros transcription factors.Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Pomalidomide-d5 is deuterium labeled Pomalidomide.Pomalidomide, the third-generationimmunomodulatory agent, acts as molecular glue.Pomalidomide interacts with the E3 ligase cereblonand induces degradation of essential Ikarostranscription factors.Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
POMHEX		Pomolic acid	
	Cat. No.: HY-131904	(Randialic acid A)	Cat. No.: HY-N6601
POMHEX, a racemic mixture and a cell-permeable pivaloyloxymethyl (POM) prodrug of HEX, is a potent, <b>ENO2</b> -specific inhibitor of enolase.		Randialic acid A (Pomolic acid) is a pentacyclic triterpene isolated from Euscaphis japonica (Tunb.). Randialic acid A (Pomolic acid) inhibits tumor cells growth and induces cell <b>apoptosis</b> .	
Purity:99.77%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg		Purity:98.14%Clinical Data:No Development ReportedSize:5 mg, 10 mg	
Poncirin		Ponicidin	
Foncian	Cat. No.: HY-N2258	(Rubescensine B)	Cat. No.: HY-N1535
Poncirin is isolated from Poncirus trifoliata with <b>anti-inflammory</b> activites. Poncirin significantly reduces mechanical hyperalgesia and allodynia in Complete Freund's Adjuvant (CFA)-induced inflammatory pain models. <b>Purity:</b> 99.55% Clinical Data: No Development Reported		Ponicidin (Rubescensine B) is a diterpenoid derived from Rabdosia rubescens, and exhibits immunoregulatory, anti-inflammatory, anti-viral and anti-cancer activity. Purity: 99.82% Clinical Pate: No Development Reported	
Size: 5 mg, 10 mg		Size: 5 mg, 10 mg	
PP1		PP121	
(AGL 1872; EI 275)	Cat. No.: HY-13804		Cat. No.: HY-10372
PP1 is a potent, and <b>Src</b> family-selective tyrosine kinase inhibitor with $IC_{s0}$ of 5 and 6 nM for Lck and Fyn, respectively.	NH2 N	PP121 is a multi-targeted kinase inhibitor with IC ₅₀ S of 10, 60, 12, 14, 2 nM for mTOR, DNK-PK, VEGFR2, Src, PDGFR, respectively.	
Purity:         98.62%           Clinical Data:         Phase 3           Size:         10 mM × 1 mL, 10 mg, 50 mg, 100 mg	" X	Purity:         99.08%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 10 mg, 50 mg, 100 mg	$\diamond$
PQ401		PR-619	
	Cat. No.: HY-13686		Cat. No.: HY-13814
PQ401 is a potent inhibitor of <b>IGF-IR</b> signaling. PQ401 inhibits IGF-I-stimulated IGF-IR autophosphorylation with an IC ₅₀ of 12.0 $\mu$ M in a series of studies in MCF-7 cells. PQ401 is effective at inhibiting IGF-I-stimulated growth of MCF-7 cells (IC ₅₀ , 6 $\mu$ M).		PR-619 is a broad-range and reversible <b>DUB</b> inhibitor with $EC_{so}^{5}$ of 3.93, 4.9, 6.86, 7.2, and 8.61 $\mu$ M for <b>USP4</b> , <b>USP8</b> , <b>USP7</b> , <b>USP2</b> , and <b>USP5</b> , respectively. PR-619 induces ER Stress and ER-Stress related apoptosis.	
Purity:       99.88%         Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 10 mg, 50 mg	~~~N. ~	Purity:       98.89%         Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 10 mg, 50 mg	5 <b>4</b> .55 57 57 72

PR-924		Pracinostat	
	Cat. No.: HY-123587	(SB939)	Cat. No.: HY-13322
PR-924 is a selective tripeptide epoxyketone immunoproteasome subunit LMP-7 inhibitor with an IC ₅₀ of 22 nM. PR-924 covalently modifies proteasomal N-terminal threonine active sites.         PR-924 inhibits growth and triggers apoptosis in multiple myeloma (MM) cells.         Purity:       >98%         Clinical Data:       No Development Reported         Size:       5 mg, 10 mg	City of the second seco	Pracinostat is a potent histone deacetylase (HDAC) inhibitor, with IC ₅₀ s of 40-140 nM, used for cancer research.         Purity:       99.82%         Clinical Data:       Phase 3         Size:       10 mM × 1 mL, 5 mg, 10 mg, 50 mg	HORIZIN
Pracinostat-d7	Cat. No.: HY-13322S	Pralatrexate	<b>Cat. No.:</b> HY-10446
Pracinostat-d7 is the deuterium labeled Pracinostat. Pracinostat is a potent <b>histone</b> <b>deacetylase (HDAC)</b> inhibitor, with <b>IC</b> ₅₀ s of 40-140 nM, used for cancer research.	HO H	Pralatrexate is an antifolate and is a potent dihydrofolate reductasean (DHFR) inhibitor with a K ₁ of 13.4 pM. Pralatrexate is a substrate for folylpolyglutamate synthetase with improved cellular uptake and retention. Purity: 99.23%	N NH2 O OKOH
Clinical Data: No Development Reported		Clinical Data: Launched	
Size: 1 mg, 10 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
Pranoprofen	<b>Cat. No.:</b> HY-B0336	Prednisone (Dehydrocortisone)	<b>Cat. No.:</b> HY-B0214
Pranoprofen is a non-steroidal anti-inflammatory agent (NSAID) for the research of keratitis or other ophthalmology diseases. Pranoprofen inhibit COX-1 and COX-2 enzymes, thus blocking arachidonic acid converted to eicosanoids and reducing prostaglandins synthesis. Purity: 99.37% Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg, 500 mg	Судерание он	Prednisone (Adasone) is a synthetic corticosteroid agent that is particularly effective as an immunosuppressant compound. Target: Others Prednisone is a synthetic corticosteroid drug that is particularly effective as an immunosuppressant drug.Purity:99.82% Clinical Data: Launched Size:10 mM × 1 mL, 500 mg, 1 g, 5 g	
Prednisone-d8 (Dehydrocortisone-d8)	<b>Cat. No.:</b> HY-B0214S	Prexasertib (LY2606368)	<b>Cat. No.:</b> HY-18174
Prednisone-d8 (Dehydrocortisone-d8) is the deuterium labeled Prednisone. Prednisone (Adasone) is a synthetic corticosteroid agent that is particularly effective as an immunosuppressant compound. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Prexasertib (LY2606368) is a selective,         ATP-competitive second-generation checkpoint kinase         1 (CHK1) inhibitor with a K ₁ of 0.9 nM and an         IC ₅₀ of <1 nM. Prexasertib inhibits CHK2 (IC ₅₀ =8 nM) and RSK1 (IC ₅₀ =9 nM).         Purity:       98.03%         Clinical Data:       Phase 2         Size:       10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	N N N N N N N N
Prexasertib dihydrochloride	Cot No : UV 101744	Prexasertib dimesylate	Cot No - UV 191745
Prexasertib dihydrochloride (LY2606368 dihydrochloride) is a selective, ATP-competitive second-generation checkpoint kinase 1 (CHK1) inhibitor with a K ₁ of 0.9 nM and an IC ₅₀ of <1 nM. Prexasertib dihydrochloride inhibits CHK2 (IC ₅₀ =8 nM) and RSK1 (IC ₅₀ =9 nM).Purity:99.41% Clinical Data:Phase 2 Size:5 mg, 10 mg, 25 mg, 50 mg, 100 mg		Prexasertib dimesylate (LY2606368 dimesylate) is a selective, ATP-competitive second-generation         checkpoint kinase 1 (CHK1) inhibitor with a K _i of         0.9 nM and an IC _{so} of <1 nM. Prexasertib	(at. NO. HT-161/4E)



Proparacaine Hydrochloride		Propylparaben	
(Proxymetacaine Hydrochloride)	Cat. No.: HY-66012	(Propyl parahydroxybenzoate; Propyl 4-hydroxybenzoate)	Cat. No.: HY-N2026
Proparacaine Hydrochloride (Proxymetacaine Hydrochloride) is a derivative of lidocaine (HY-B0185), with immunomodulatory effect and glucocorticomimetic activity.	H,N, Jown	Propylparaben (Propyl parahydroxybenzoate) is an antimicrobial preservative which can be produced naturally by plants and bacteria. Propylparaben is prevalently used in cosmetics, pharmaceuticals, and foods.	HO
Purity:         99.76%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 100 mg		Purity:98.93%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 100 mg, 1 g	
Propyl paraben sodium (Propyl parahydroxybenzoate Propyl 4-hydroxybenzoate sodium)	cat. No.: HY-N2026A	Prosapogenin A (Progenin III)	Cat. No.: HY-N6940
Propylparaben sodium (Propyl parahydroxybenzoate) is an antimicrobial preservative which can be produced naturally by plants and bacteria. Propylparaben sodium is prevalently used in cosmetics, pharmaceuticals, and foods.	NaO	Prosapogenin A, a natural product from Veratrum, induces apoptosis in human cancer cells in vitro via inhibition of the STAT3 signaling pathway and glycolysis.	A Contraction of the second se
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg		Purity:99.55%Clinical Data:No Development ReportedSize:5 mg, 10 mg	
DPOTAC ELT-3 degrader 1			
	Cat No: HY-114323	FROTAC-0412	Cat No: HY-141881
PROTAC FLT-3 degrader 1 is a <b>von</b> <b>Hippel-Lindau</b> -based PROTAC <b>FLT-3</b> internal tandem duplication (ITD) degrader with an IC _{s0} 0.6 nM. Anti-proliferative activity; apoptosis induction.	jaturnozounat	PROTAC-O4I2 is a <b>PROTAC</b> targets splicing factor 3B1 ( <b>SF3B1</b> ). PROTAC-O4I2 induces FLAG-SF3B1 degradation with an IC _{s0} value of 0.244 $\mu$ M in K562 cells. PROTAC-O4I2 also induces cellular <b>apoptosis</b> in K562 WT cells.	
Purity:98.70%Clinical Data:No Development ReportedSize:1 mg, 5 mg, 10 mg		Purity:98.00%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	o'
Protosannanin B		DS_11//5	
((-)-Protosappanin B)	Cat No HY-N0800	L 2-TT+2	Cat No : HY-18008
Protosappanin B is a phenolic compound extracted from Lignum Sappan. Anti-cancer activity.         Protosappanin B induces apoptosis and causes G1 cell cycle arrest in human bladder cancer cells.         Purity:       99.46%         Clinical Data:       No Development Reported	HO HO HO HO	PS-1145 is an IkB kinase (IKK) inhibitor with an IC ₅₀ of 88 nM. Purity: 99.88% Clinical Data: No Development Reported	NO CHART
Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	0 mg
PSB 0474	<b>Cat. No.:</b> HY-108654	Pseudolaric Acid B	<b>Cat. No.:</b> HY-N6939
PSB 0474 (3-phenacyl-UDP) is a selective and potent <b>P2Y</b> ₆ receptor agonist with an <b>EC</b> _{so} of 70 nM. PSB 0474 inhibits cell proliferation, increases NO release in astrocytes and microglia cells. PSB 0474 induces astrocytes <b>apoptosis</b> .		Pseudolaric Acid B is a diterpene isolated from the root of Pseudolarix kaempferi Gorden (pinaceae), has anti-cancer, antifungal, and antifertile activities, and shows immunosuppressive activity on T lymphocytes.	
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:         99.47%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	

Psoralen		P1C-028	<b>6</b> • • • • • • • • • • • • • • • • • • •
(Ficusin)	Cat. No.: HY-N0053		Cat. No.: HY-103696
Psoralen (Ficusin) is a coumarin isolated from the seeds of Fructus Psoraleae. Psoralen exhibits a wide range of biological properties, including anti-cancer, antioxidant, antidepressant, anticancer, antibacterial, and antiviral, et al.	°	PTC-028 is an orally bioavailable inhibitor of stem cell factor <b>BMI-1</b> in ovarian cancer. PTC-028 selectively inhibits cancer cells whereas normal cells remain unaffected. PTC-028 downregulates BMI-1, inducing caspase-mediated apportosis	
Purity:         99.92%           Clinical Data:         Phase 1           Size:         10 mM × 1 mL, 10 mg, 50 mg, 100 mg		Purity:       ≥98.0%         Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	)0 mg
PU02	Cat. No.: HY-103118	Puerarin 6''-O-Xyloside	<b>Cat. No.:</b> HY-N2135
PU02, a derivative of 6-MP (HY-13677), is a negative allosteric modulator (NAM) of $5-HT_3$ receptor, with IC _{so} values of 0.36 and 0.73 $\mu$ M in HEK293 cells transfected with human 5-HT ₃ A and 5-HT ₃ AB receptors respectively.	HZ N S	Puerarin 6"-O-Xyloside, isolated from radix of Pueraria lobata (Willd.), possesses snti-osteoporotic and anti-tumor activity. Puerarin 6"-O-Xyloside induces the mitochondria-mediated apoptosis pathway.	HO, CO CHARACH
Purity:99.29%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg, 50 mg		Purity:99.52%Clinical Data:No Development ReportedSize:1 mg, 5 mg	1076.
Pulsatilla saponin D		Puromycin aminonucleoside	
Pulsatilla saponin D (SB365), isolated from the root of Pulsatilla koreana Nakai, is an anti-tumor agent.		Puromycin aminonucleoside (NSC 3056) is the aminonucleoside portion of the antibiotic puromycin, and used in nephrosis animal models. Puromycin aminonucleoside induces <b>apoptosis</b> .	
Purity:98.47%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg	HOLO .	Purity:99.67%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 5	H ₂ N 500 mg, 1 g
		Pygenic acid A	
(NG-60)	Cat No: HY-18299A	rygenic acid A	Cat No: HY-N1823
Purvalanol A is a potent <b>CDK</b> inhibitor, which inhibits cdc2-cyclin B, cdk2-cyclin A, cdk2-cyclin E, cdk4-cyclin D1, and cdk5-p35 with <b>IC</b> _{so} s of 4, 70, 35, 850, 75 nM, resepctively.		Pygenic acid A is a natural compound that can be found in Prunella vulgaris. Pygenic acid A induces <b>apoptosis</b> in metastatic breast cancer cells. Pygenic acid A can be used for the research of diabetes, inflammatory diseases, and cancers.	
Purity:         99.11%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 10 mg, 25 mg, 50 mg, 100 mg		Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg	, , , , , , , , , , , , , , , , , , ,
Pyoluteorin	<b>Cat. No.:</b> HY-114979	PYR-41	<b>Cat. No.:</b> HY-13296
Pyoluteorin is an <b>antibiotic</b> that inhibits Oomycete fungi, including the plant pathogen Pythium ultimum, and suppresses plant diseases caused by this fungus. Pyoluteorin induces human triple-negative breast cancer MDA-MB-231 cells <b>apoptosis</b> in vitro.		PYR-41 is a selective and cell permeable inhibitor of ubiquitin-activating enzyme E1 with an $IC_{s0}$ of < 10 $\mu$ M, with little activity at E2 and E3.	
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg		Purity:     ≥ 98.0%       Clinical Data:     No Development Reported       Size:     10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	

Pyrazoloacridine           (NSC 366140; PD 115934)         Cat. No.: HY-108969	Pyrogallol	<b>Cat. No.</b> : HY-N1579
Pyrazoloacridine (NSC 366140), an intercalating agent with anti-cancer activity, inhibits the activity of <b>topoisomerases</b> 1 and 2. Pyrazoloacridine (NSC 366140) exhibits an <b>IC</b> ₅₀ of 1.25 $\mu$ M in K562 myeloid leukemia cells for 24 h treatment.	Pyrogallol is a polyphenol compound, which has anti-fungal and anti-psoriatic properties. Pyrogallol is a reductant that is able to generate free radicals, in particular superoxide anions.	OH OH
Purity:     ≥ 98.0%     O ^{ch}	Purity:99.98%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 500 mg, 1 g	🔨 ОН
Pyroxamide Cat. No.: HY-13216	QTX125	<b>Cat. No.</b> : HY-120448
Pyroxamide is a potent inhibitor of <b>histone</b> <b>deacetylase 1 (HDAC1)</b> with an <b>ID</b> ₅₀ of 100 nM. Pyroxamide can induce apoptosis and cell cycle arrest in leukemia.	QTX125 is a potent and highly selective <b>HDAC6</b> inhibitor. QTX125 exhibits excellent selectivity over other HDACs. QTX125 has antitumor effects.	10-0-4-14-0-14 of
Purity:         99.73%           Clinical Data:         Phase 1           Size:         10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg	Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
QTX125 TFA Cat. No.: HY-120448A	Quercetin	<b>Cat. No.:</b> HY-18085
QTX125 TFA is a potent and highly selective <b>HDAC6</b> inhibitor. QTX125 TFA exhibits excellent selectivity over other HDACs. QTX125 has antitumor effects.	Quercetin, a natural flavonoid, is a stimulator of recombinant SIRT1 and also a PI3K inhibitor with IC ₅₀ of 2.4 $\mu$ M, 3.0 $\mu$ M and 5.4 $\mu$ M for PI3K $\gamma$ , PI3K $\delta$ and PI3K $\beta$ , respectively.	но странон
Purity:     >98%       Clinical Data:     No Development Reported       Size:     5 mg, 10 mg, 25 mg, 50 mg, 100 mg	Purity:         98.02%           Clinical Data:         Phase 4           Size:         10 mM × 1 mL, 500 mg, 1 g, 5 g	OH O
Quercetin dihydrate Cat. No.: HY-N0146	Quercetin-d3	<b>Cat. No.</b> : HY-18085S1
Quercetin dihydrate, a natural flavonoid, is a stimulator of recombinant SIRT1 and a PI3K inhibitor with IC ₅₀ s of 2.4 $\mu$ M, 3.0 $\mu$ M and 5.4 $\mu$ M for PI3K $\gamma$ , PI3K $\delta$ and PI3K $\beta$ , respectively.	Quercetin-d3 is the deuterium labeled Quercetin. Quercetin, a natural flavonoid, is a stimulator of recombinant SIRT1 and also a PI3K inhibitor with IC ₅₀ of 2.4 $\mu$ M, 3.0 $\mu$ M and 5.4 $\mu$ M for PI3K $\gamma$ , PI3K $\delta$ and PI3K $\beta$ , respectively.	он о но страна в страна в страна
Purity:         ≥96.0%           Clinical Data:         Phase 4           Size:         10 mM × 1 mL, 500 mg	Purity:>98%Clinical Data:No Development ReportedSize:2.5 mg, 25 mg	
Quercetin-d5 Cat. No.: HY-180855	Quiflapon (MK-591)	<b>Cat. No.:</b> HY-10037
Quercetin-d5 is a deuterium labeled Quercetin. Quercetin, a natural flavonoid, is a stimulator of recombinant SIRT1 and also a PI3K inhibitor with IC ₅₀ of 2.4 $\mu$ M, 3.0 $\mu$ M and 5.4 $\mu$ M for PI3K $\gamma$ , PI3K $\delta$ and PI3K $\beta$ , respectively.	Quiflapon (MK-591) is a selective and specific <b>5-lipoxygenase-activating protein (FLAP)</b> inhibitor with an $IC_{s0}$ of 1.6 nM in a FLAP binding assay.	Ctro Ctro
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	Purity:         99.44%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	

Quiflapon sodium		Quinacrine dihydrochloride	
(MK-591 sodium)	Cat. No.: HY-50714	(Mepacrine dihydrochloride; SN-390 dihydrochloride)	Cat. No.: HY-13735A
Quiflapon sodium (MK-591 sodium) is a selective and specific 5-Lipoxygenase-activating protein (FLAP) inhibitor. Quiflapon sodium is an orally active Leukotriene biosynthesis inhibitor. Induces apoptosis.	Carlos Carlos	Quinacrine (Mepacrine) dihydrochloride is an orally bioavailable <b>antimalaria</b> l agent, which possess anticancer effect both in vitro and vivo. Quinacrine dihydrochloride suppresses NF-kB and activate p53 signaling, which results in the induction of the <b>apoptosis</b> .	HN HCI
Purity: 98.65%		Purity: 99.01%	
Clinical Data: No Development Reported		Clinical Data: Phase 2	City 1999 with Sector
Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Size: 10 mM × 1 mL, 100 mg, 500 mg	
Quinacrine hydrochloride hydrate (Mepacrine hydr	ochloride	Quisinostat	
hydrate; SN-390 hydrochloride hydrate)	Cat. No.: HY-13735B	(JNJ-26481585)	Cat. No.: HY-15433
Quinacrine hydrochloride hydrate (Mepacrine hydrochloride hydrate) is an antimalarial agent, which possess anticancer effect both in vitro and vivo. Quinacrine hydrochloride hydrate suppresses NF-kB and activates p53 signaling, which results in the induction of the apoptosis.Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	HO HO	Quisinostat (JNJ-26481585) is a potent,         second-generation and orally active pan-HDAC         inhibitor (HDACi), with IC ₅₀ values ranging from         0.11 nM to 0.64 nM for HDAC1, HDAC2, HDAC4, HDAC10         and HDAC11. Quisinostat has a broad spectrum         antitumoral activity.         Purity:       98.02%         Clinical Data:       Phase 2         Size:       10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	after NN H (N)
Quisinostat dihydrochloride		Quizartinib	
(JNJ-26481585 dihydrochloride)	Cat. No.: HY-15433A	(AC220)	Cat. No.: HY-13001
Quisinostat dihydrochloride (JNJ-26481585 dihydrochloride) is an orally available, potent pan- <b>HDAC</b> inhibitor with IC ₅₀ s of 0.11 nM, 0.33 nM, 0.64 nM, 0.46 nM, and 0.37 nM for HDAC1, HDAC2, HDAC4, HDAC10 and HDAC11, respectively.	N N N H CI H-CI	Quizartinib (AC220) is an orally active, highly selective and potent second-generation type II <b>FLT3 tyrosine kinase</b> inhibitor, with a K _d of 1.6 nM. Quizartinib inhibits wild-type FLT3 and FLT3-ITD autophosphorylation in MV4-11 cells with IC ₅₀ s of 4.2 and 1.1 nM, respectively.	andraget
Purity: >98%		Purity: 99.01%	
Clinical Data: No Development Reported		Clinical Data: Launched	
Size: 1 mg, 5 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 2	00 mg
R1530		R406	
	Cat. No.: HY-13737		Cat. No.: HY-12067
R1530 is a highly potent, orally active, dual-acting <b>mitosis/angiogenesis</b> inhibitor, with anti-tumor and anti-angiogenic activities. R1530 is a multikinase inhibitor which binds to 31 kinases with K _a s of <500 nM.		R406 is an orally available and competitive <b>Syk/FLT3</b> inhibitor for ATP binding with a K ₁ of 30 nM, potently inhibits Syk kinase activity in vitro with an IC ₅₀ of 41 nM, measured at an ATP concentration corresponding to its K _m value.	giriat q.
Purity: 99.06% Clinical Data: No Development Reported		Purity: 96.67% Clinical Data: No Development Reported	
Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
R406 free base		R547	
	Cat. No.: HY-11108		Cat. No.: HY-10014
R406 free base is an orally available and competitive <b>Syk/FLT3</b> inhibitor for ATP binding with a $K_i$ of 30 nM, potently inhibits Syk kinase activity in vitro with an IC _{s0} of 41 nM, measured at an ATP concentration corresponding to its $K_m$ value.	Jog and and	R547 is a potent, selective and orally active ATP-competitive CDK inhibitor, with K ₅ of 2 nM, 3 nM and 1 nM for CDK1/cyclin B, CDK2/cyclin E and CDK4/cyclin D1, respectively.	
Purity: 99.69%		Purity: 99.66%	o=s=o
Clinical Data: No Development Reported		Clinical Data: No Development Reported	
SIZE: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Size: 10 mM × 1 mL, 10 mg, 50 mg	

RA-9		RA375	
	Cat. No.: HY-136528	10.075	Cat. No.: HY-136563
RA-9 is a potent and selective proteasome-associated <b>deubiquitinating enzymes</b> ( <b>DUBs</b> ) inhibitor with favorable toxicity profile and anticancer activity.	° gran franko	RA375 is a <b>RPN13 (26S proteasome regulatory</b> <b>subunit)</b> inhibitor. RA375 activates UPR signaling, ROS production and apoptosis. RA375 exhibits ten-fold greater activity against cancer lines than RA190, reflecting its nitro ring substituents and the addition of a chloroacetamide warhead	o y C y C y C y C y C y C y C y C y C y
Purity:         98.12%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity:       >98%         Clinical Data:       No Development Reported         Size:       5 mg, 10 mg, 50 mg, 100 mg	à
Rabeprazole (LY307640)	<b>Cat. No.:</b> HY-B0656	Rabeprazole sodium (LY307640 sodium)	<b>Cat. No.</b> : HY-B0656A
Rabeprazole (LY307640) is a second-generation proton <b>pump inhibitor</b> ( <b>PPI</b> ) that irreversibly inactivates gastric H ⁺ /K ⁺ -ATPase. Rabeprazole induces <b>apoptosis</b> . Rabeprazole acts as an uridine nucleoside ribohydrolase ( <b>UNH</b> ) inhibitor with an <b>IC</b> _{s0} of 0.3 $\mu$ M.	C the source of the second sec	Rabeprazole sodium (LY307640 sodium) is a second-generation proton <b>pump inhibitor (PPI)</b> that irreversibly inactivates gastric H ⁺ /K ⁺ -ATPase. Rabeprazole sodium induces <b>apoptosis</b> .	Q N
Purity:     >98%       Clinical Data:     Launched       Size:     1 mg, 5 mg		Purity:       99.17%         Clinical Data:       Launched         Size:       10 mM × 1 mL, 10 mg, 50 mg	
Rabeprazole-d3 sodium		Rabeprazole-d4	
(LY307640-d3 sodium)	Cat. No.: HY-B0656AS1	(LY307640-d4)	Cat. No.: HY-B0656S
Rabeprazole-d3 (LY307640-d3) sodiumis the deuterium labeled Rabeprazole sodium. Rabeprazole sodium (LY307640 sodium) is a second-generation proton <b>pump inhibitor</b> ( <b>PPI</b> ) that irreversibly inactivates gastric H ⁺ /K ⁺ -ATPase. Rabeprazole sodium induces <b>apoptosis</b> .	Chi Harris	Rabeprazole D4 (LY307640 D4) is a deuterium labeled Rabeprazole. Rabeprazole is a second-generation proton <b>pump inhibitor</b> ( <b>PPI</b> ) that irreversibly inactivates gastric H*/K*-ATPase. Rabeprazole induces <b>apoptosis</b> .	
Clinical Data:         No Development Reported           Size:         1 mg, 5 mg		Clinical Data: No Development Reported Size: 1 mg, 5 mg	
Rabeprazole-d4 sodium	Cat No . HY POSESAS	(B02)	Cat No : HV 101462
Rabeprazole-d4 sodium (LY307640-d4 sodium) is the deuterium labeled Rabeprazole sodium. Rabeprazole sodium (LY307640 sodium) is a second-generation proton <b>pump inhibitor</b> ( <b>PPI</b> ) that irreversibly inactivates gastric H*/K*-ATPase. Rabeprazole sodium induces <b>apoptosis</b> .		RAD51 Inhibitor B02 (B02) is an inhibitor of human RAD51 with an IC $_{\rm so}$ of 27.4 $\mu M.$	
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:         99.87%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	Ö 10 mg
Raddeanin A	<b>Cat. No.:</b> HY-N0819	RAF265 (CHIR-265)	<b>Cat. No.</b> : HY-10248
Raddeanin A is a natural triterpenoid saponin component of Anemone raddeana, with anti-cancer activities. Raddeanin A exerts anticancer effect on human osteosarcoma via the ROS/JNK and NF-kB signal pathway.	$(\mathcal{A}_{\mathcal{A}}^{(n)}) = (\mathcal{A}_{\mathcal{A}}^{(n)}) $	RAF265 is a potent <b>RAF/VEGFR2</b> inhibitor.	Storapt
Purity:98.15%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 20 mg	κο ^{-γ} α	Purity:         99.98%           Clinical Data:         Phase 2           Size:         10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg	



RET-IN-11	<b>Cat. No.:</b> HY-144131	Reveromycin A	<b>Cat. No.:</b> HY-129337
RET-IN-11 is a potent and selective RET inhibitor         with IC ₅₀ s of 6.20 nM, 18.68 nM for RET and         RETY804M, respectively. RET-IN-11 shows         anti-proliferation and migration activity in         CCDC6-RET-driven LC-2/ad cells. RET-IN-11 induces         cell apoptosis.         Purity:       >98%         Clinical Data:       No Development Reported         Size:       1 mg, 5 mg		Reveromycin A, a benzoquinoid antibiotic isolated from the genus Streptomyces, is a selective inhibitor of protein synthesis in eukaryotic cells. Reveromycin A inhibits bone resorption by inducing apoptosis specifically in osteoclasts.Purity:>98% Clinical Data: Size:No Development Reported Size:1 mg, 5 mg, 10 mg, 25 mg	and a set of the set o
RGD peptide (GRGDNP)	<b>Cat. No.:</b> HY-P1740	RGD peptide (GRGDNP) (TFA)	<b>Cat. No</b> .: HY-P1740A
RGD peptide (GRGDNP) acts as an inhibitor of integrin-ligand interactions and plays an important role in cell adhesion, migration, growth, and differentiation.		RGD peptide (GRGDNP) (TFA) acts as an inhibitor of integrin-ligand interactions and plays an important role in cell adhesion, migration, growth, and differentiation.	
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	H ₀ N  NH	Purity:99.25%Clinical Data:No Development ReportedSize:1 mg, 5 mg	<i>इ</i> र्गू 04
Rhapontin		Rhosin	
(Rhaponiticin)	Cat. No.: HY-N0671		Cat. No.: HY-12646A
Rhapontin (Rhaponiticin), a component of rhubarb (Rheum officinale Baillon), induces <b>apoptosis</b> resulting in suppression of proliferation of human stomach cancer KATO III cells.	HO, CH, CH, CH, CH, CH, CH, CH, CH, CH, CH	Rhosin is a potent, specific RhoA subfamily <b>Rho</b> <b>GTPases</b> inhibitor, which specifically binds to RhoA to inhibit RhoA-GEF interaction with a $K_d$ of ~ 0.4 uM, and does not interact with Cdc42 or Rac1, nor the GEF, LARG. Rhosin induces cell <b>apoptosis</b> .	Que Land
Purity:99.67%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 20 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
Rhosin hydrochloride		RHPS4	
	Cat. No.: HY-12646		Cat. No.: HY-101089
Rhosin hydrochloride is a potent, specific RhoA subfamily <b>Rho GTPases</b> inhibitor. Rhosin hydrochloride specifically binds to RhoA to inhibit RhoA-GEF interaction with a $K_d$ of ~ 0.4 uM, and does not interact with Cdc42 or Rac1, nor the GEF, LARG.	Contraction of the second seco	RHPS4 is a potent <b>telomerase</b> inhibitor (IC ₅₀ = 0.33 $\mu$ M). RHPS4 is a DNA damage inducer.	- CHA - 250
Purity:99.93%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg		Purity:98.62%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg	
Picolinostat		Pigocartib	
(ACY-1215; Rocilinostat)	<b>Cat. No.:</b> HY-16026	(ON-01910)	Cat. No.: HY-12037A
Ricolinostat (ACY-1215) is a potent and selective HDAC6 inhibitor, with an $IC_{s0}$ of 5 nM. ACY-1215 also inhibits HDAC1, HDAC2, and HDAC3 with $IC_{s0}$ s of 58, 48, and 51 nM, respectively.	Q [*] [*] [*] [*] [*] [*]	Rigosertib (ON-01910) is a multi-kinase inhibitor and a selective anti-cancer agent, which induces apoptosis by inhibition the <b>PI3 kinase/Akt</b> pathway, promots the phosphorylation of histone H2AX and induces G2/M arrest in cell cycle.	, С.С. С.С. К.С.
Purity:         99.83%           Clinical Data:         Phase 2           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity:         98.81%           Clinical Data:         Phase 3           Size:         5 mg, 10 mg, 50 mg, 100 mg	

Rigosertib sodium		Rilmenidine	
(ON-01910 sodium)         Rigosertib sodium (ON-01910 sodium) is a multi-kinase inhibitor and a selective anti-cancer agent, which induces apoptosis by inhibition the         PI3K/Akt pathway, promotes the phosphorylation of histone H2AX and induces G2/M arrest in cell cycle.         Purity:       99.49%         Clinical Data:       Phase 3         Size:       10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	Cat. No.: HY-12037	Rilmenidine, an innovative antihypertensive agent, is an orally active, selective II imidazoline receptor agonist. Rilmenidine is an alpha 2-adrenoceptor agonist. Rilmenidine induces autophagy.         Purity:       >98%         Clinical Data:       Launched         Size:       1 mg, 5 mg	Cat. No.: HY-100490
Rilmenidine hemifumarate	<b>Cat. No.:</b> HY-100490A	Rilmenidine phosphate	<b>Cat. No.:</b> HY-100490B
Rilmenidine hemifumarate, an innovative antihypertensive agent, is an orally active, selective I1 imidazoline receptor agonist.         Rilmenidine hemifumarate is an alpha         2-adrenoceptor agonist.         Rilmenidine hemifumarate is an alpha         2-adrenoceptor agonist.         Rilmenidine hemifumarate is an alpha         2-adrenoceptor agonist.         Rilmenidine hemifumarate         induces autophagy.         Purity:       99.82%         Clinical Data:       Launched         Size:       5 mg, 10 mg		Rilmenidine phosphate, an innovative         antihypertensive agent, is an orally active,         selective II imidazoline receptor agonist.         Rilmenidine phosphate is an alpha 2-adrenoceptor         agonist. Rilmenidine phosphate induces autophagy.         Purity:       ≥98.0%         Clinical Data:       Launched         Size:       5 mg, 10 mg, 25 mg	N N N H HO-P-OH OH
Rilmenidine-d4	<b>Cat. No.:</b> HY-100490S	Rimiducid (AP1903)	<b>Cat. No.:</b> HY-16046
Rilmenidine-d4 is the deuterium labeledRilmenidine. Rilmenidine, an innovativeantihypertensive agent, is an orally active,selective I1 imidazoline receptor agonist.Rilmenidine is an alpha 2-adrenoceptor agonist.Rilmenidine induces autophagy.Purity:> 98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Rimiducid (AP1903) is a dimerizer agent that acts         by cross-linking the FKBP domains. Rimiducid         (AP1903) dimerizes the Caspase 9 suicide switch         and rapidly induces apoptosis.         Purity:       99.81%         Clinical Data:       Phase 3         Size:       2 mg, 5 mg, 10 mg, 50 mg, 100 mg	ng Agantanaga Ng Ng Ng Ng Ng Ng Ng Ng Ng Ng Ng Ng Ng
RIPGBM	<b>Cat. No.:</b> HY-122910	Ripretinib (DCC-2618)	<b>Cat. No.:</b> HY-112306
RIPGBM is a selective inducer of <b>apoptosis</b> in glioblastoma multiforme (GBM) cancer stem cells (CSCs) with an $EC_{s0}$ of $\leq$ 500 nM.Purity:99.89%	O N O N O O O O O O O O O O O O O O O O	Ripretinib (DCC-2618) is an orally bioavailable, selective KIT and PDGFRA switch-control inhibitor.	CHING OF CHING
Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	00 mg	Clinical Data:         Launched           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
Ritonavir         (ABT 538; RTV)         Ritonavir (ABT 538) is an inhibitor of HIV         protease used to treat HIV infection and AIDS.         Ritonavir is also a SARS-CoV 3CL ^{pro} inhibitor         with an IC _{so} of 1.61 μM.	<b>Cat. No.:</b> HY-90001	Ritonavir-13C,d3 (ABT 538-13C,d3; RTV-13C,d3) Ritonavir-13C,d3 (ABT 538-13C,d3) is the 13C- and deuterium labeled Ritonavir. Ritonavir (ABT 538) is an inhibitor of HIV protease used to treat HIV infection and AIDS. Ritonavir is also a SARS-CoV 3CIP [®] inhibitor with an IC of 1.61 µM	Cat. No.: HY-9000151
Purity:         99.95%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 500 mg	U	Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	U

Ritonavir-d6		Riviciclib hydrochloride	C-+ N UV 10550
Ritonavir-d6 (ABT 538-d6) is the deuterium labeled Ritonavir. Ritonavir (ABT 538) is an inhibitor of HIV protease used to treat HIV infection and AIDS. Ritonavir is also a SARS-CoV 3CL ^{pro} inhibitor with an IC ₅₀ of 1.61 $\mu$ M. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg		Riviciclib hydrochloride (P276-00) is a potent         cyclin-dependent kinase (CDK) inhibitor, which         inhibits CDK9-cyclinT1, CDK4-cyclin D1, and         CDK1-cyclinB with IC ₅₀ s of 20 nM, 63 nM, and 79 nM,         respectively.         Purity:       99.01%         Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	HO HO H-CI
RKI-1447 dihydrochloride	<b>Cat. No.:</b> HY-110339	RMS3	<b>Cat. No.:</b> HY-146096
RKI 1447 dihydrochloride is a potent and selective         ROCK inhibitor with IC _{so} s of 14.5 and 6.2 nM for         ROCK1 and ROCK2, respectively. RKI 1447         dihydrochloride suppresses colorectal carcinoma         cell growth and promotes apoptosis.         Purity:       98.04%         Clinical Data:       No Development Reported         Size:       5 mg 10 mg 25 mg 50 mg 100 mg	HO THE HOLD	RMS3, a tetrandrine analogue, is a potent P-glycoprotein (P-gp) inhibitor. RMS3 has markedly antiproliferative and cytotoxic effects on cancer cells. RMS3 causes PARP cleavage, a marker for cells undergoing apoptosis. RMS3 has strong anticancer property. Purity: >98% Clinical Data: No Development Reported Size: 1 mg 5 mg	
ن ,ن ,ن		5,	
RMS5		Ro 08-2750	
	Cat. No.: HY-146097		Cat. No.: HY-108466
RMS5, a tetrandrine analogue, is a potentP-glycoprotein (P-gp) inhibitor. RMS5 has markedly antiproliferative and cytotoxic effects on cancer cells. RMS5 slightly diminishes the expression of the anti-apoptotic Bcl-2 family proteins Bcl-XL and Mcl-1.Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Ro 08-2750 is a non-peptide and reversible nervegrowth factor (NGF) inhibitor which binds to NGF,and with an ICand with an ICso of ~ 1 μM. Ro 08-2750 inhibitsNGF binding to p75NTR selectively over TRKA. Ro08-2750 is a selectiveMSI RNA-bindingactivityinhibitor, with an ICso of 2.7 μM.Purity:95.40%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 1 mg, 5 mg	OF THE NOTION
Ro 90-7501		Ro-3306	Cat No. LIV 12520
Ro 90-7501 is an <b>amyloid</b> $\beta_{42}$ (A $\beta_{42}$ ) fibril <b>assembly</b> inhibitor that reduces A $\beta_{42}$ -induced cytotoxicity (EC ₅₀ of 2 $\mu$ M). Ro 90-7501 inhibits ATM phosphorylation and DNA repair.		Ro-3306 is a potent and selective inhibitor of CDK1, with $K_1$ s of 20 nM, 35 nM and 340 nM for CDK1, CDK1/cyclin B1 and CDK2/cyclin E, respectively.	S N S
Purity:       >98%         Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	00 mg	Purity:         98.92%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	√~N
Ro24-7429	<b>Cat. No</b> .: HY-19149	ROC-325	<b>Cat. No.:</b> HY-103706
Ro24-7429 is a potent and orally active HIV-1 transactivator protein <b>Tat</b> antagonist. Ro24-7429 is also a <b>runt-related transcription factor 1</b> ( <b>RUNX1</b> ) inhibitor. Ro24-7429 has anti-HIV, antifibrotic and anti-inflammatory effects. <b>Purity:</b> 99.90%		ROC-325 is a potent and orally active <b>autophagy</b> inhibitor with a strong anticancer activity. ROC-325 induces the deacidification of lysosomes, accumulation of autophagosomes, and disrupted autophagic flux. ROC-325 also induces renal cell carcinoma <b>apoptosis</b> . <b>Purity:</b> 99.91%	
Clinical Data:       No Development Reported         Size:       5 mg, 10 mg, 25 mg, 50 mg, 100 mg	ŕ	Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	00 mg



RWJ-56110	<b>Cat. No.</b> : HY-108556	RWJ-56110 dihydrochloride	<b>Cat. No.</b> : HY-108556A
RWJ-56110 is a potent, selective, peptide-mimetic inhibitor of <b>PAR-1</b> activation and internalization (binding $IC_{50}$ =0.44 uM) and shows no effect on PAR-2, PAR-3, or PAR-4.	gangin	RWJ-56110 dihydrochloride is a potent, selective, peptide-mimetic inhibitor of <b>PAR-1</b> activation and internalization (binding $IC_{so}$ =0.44 uM) and shows no effect on PAR-2, PAR-3, or PAR-4.	a manufacture
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg		Purity:99.54%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
S-Allyl-L-cysteine	Cat. No.: HY-W013573	S-Allylmercaptocysteine	<b>Cat. No.</b> : HY-145532
S-Allyl-L-cysteine, one of the organosulfur compounds found in AGE, possess various biological effects including neurotrophic activity, anti-cancer activity, anti-inflammatory activity.	S C OH	S-allylmercaptocysteine, an organic sulfur compound extracted from garlic, has anti-inflammatory and anti-oxidative effects for various pulmonary diseases.	S-S-NH2 OH
Purity:         98.64%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 10 mg, 25 mg, 50 mg, 100 mg		Purity:≥95.0%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
S-Trityl-L-cysteine (NSC 83265; S-Tritylcysteine; 3-Tritylthio-L-alanine)	Cat. No.: HY-W011102	S116836	<b>Cat. No.:</b> HY-123450
S-Trityl-L-cysteine (NSC 83265) is a selective and allosteric <b>kinesin Eg5</b> inhibitor with an $IC_{50}$ of 1 $\mu$ M for the inhibition of basal ATPase activity and 140 nM for the microtubule-activated ATPase activity. S-Trityl-L-cysteine has antitumor activities.		S116836, a potent, orally active <b>BCR-ABL</b> tyrosine kinase inhibitor, blocks both wild-type as well as T315I Bcr-Abl.	Jun Cto
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg		Purity:99.60%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
S130	<b>Cat. No.:</b> HY-112818	S2116	<b>Cat. No.</b> : HY-136522
S130 is a high affinity, selective inhibitor of ATG4B (a major cysteine protease) with an $IC_{s0}$ of 3.24 $\mu$ M. S130 suppresses autophagy flux.	C the second sec	S2116, a N-alkylated tranylcypromine (TCP) derivative, is a potent <b>lysine-specific demethylase</b> <b>1 (LSD1)</b> inhibitor. S2116 increases H3K9 methylation and reciprocal H3K27 deacetylation at super-enhancer regions.	
Purity:99.31%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
S2157	<b>Cat. No.:</b> HY-136523	S65487 (VOB560)	<b>Cat. No.</b> : HY-138697
S2157, a N-alkylated tranylcypromine (TCP) derivative, is a potent <b>lysine-specific demethylase</b> <b>1 (LSD1)</b> inhibitor. S2157 increases H3K9 methylation and reciprocal H3K27 deacetylation at super-enhancer regions.	Relative stereochemistry	S65487 (VOB560), a potent and selective <b>BCL-2</b> inhibitor, is a prodrug of S55746. S65487 is also active on BCL-2 mutations, such as G101V and D103Y. S65487 has poor affinity with MCL-1, BFL-1 and BCL-XL. S65487 induces <b>apoptosis</b> and has anticaner activities.	
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg		Purity:         99.10%           Clinical Data:         Phase 2           Size:         5 mg, 10 mg, 25 mg, 50 mg, 100 mg	~~~ ~ "

S65487 hydrochloride		S65487 sulfate	
(VOB560 hydrochloride)	Cat. No.: HY-138697B	(VOB560 sulfate)	Cat. No.: HY-138697A
S65487 (VOB560) hydrochloride, a potent and selective Bcl-2 inhibitor, is a prodrug of S55746.S65487 hydrochloride is also active on BCL-2 mutations, such as G101V and D103Y. S65487 hydrochloride has poor affinity with MCL-1, BFL-1 and BCL-XL.Purity:99.67% Clinical Data:Size:5 mg, 10 mg, 25 mg, 50 mg, 100 mg		S65487 (VOB560) sulfate, a potent and selectiveBcl-2 inhibitor, is a prodrug of S55746. S65487sulfate is also active on BCL-2 mutations, such asG101V and D103Y. S65487 sulfate has poor affinitywith MCL-1, BFL-1 and BCL-XL. S65487 sulfateinduces apoptosis and has anticaner activities.Purity:98.08%Clinical Data:Phase 2Size:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	ad the second
Sabizabulin (VERU-111; ABI-231)	<b>Cat. No.</b> : HY-120599	Sacubitril/Valsartan (LCZ696)	<b>Cat. No.</b> : HY-18204A
VERU-111 (ABI-231) is a potent and orally active $\alpha$ and $\beta$ tubulin inhibitor, which displays strong antiproliferative activity, with an average IC ₅₀ of 5.2 nM against panels of melanoma and prostate cancer cell lines. Purity: 98.02% Clinical Data: Phase 3	$(\mathbf{y}_{\mathbf{y}_{n}}^{H})_{\mathbf{y}_{n}} (\mathbf{y}_{n})_{\mathbf{y}_{n}} (\mathbf{y}_{n})_{\mathbf{y}} (\mathbf{y}_{n})_{\mathbf{y}_{n}} (\mathbf{y}_{n})$	Sacubitril/Valsartan (LCZ696), comprised Valsartan and Sacubitril (AHU377) in 1:1 molar ratio, is a first-in-class, orally bioavailable, and dual-acting <b>angiotensin receptor-neprilysin (ARN)</b> inhibitor for hypertension and heart failure. <b>Purity:</b> 99.99% <b>Clinical Data:</b> Launched	
Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg	
Sal003	<b>Cat. No.</b> : HY-15969	Salermide	<b>Cat. No.:</b> HY-101073
Sal003 is a potent, specific and cell-permeable inhibitor of the <b>eukaryotic translation initiation</b> <b>factor 2$\alpha$ (eIF2$\alpha$) phosphatase</b> . Sal003 is a derivative of salubrinal.		Salermide is an inhibitor of <b>Sirt1</b> and <b>Sirt2</b> ; can cause strong cancer-specific apoptotic cell death.	COH
Purity:99.75%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg		Purity:       ≥98.0%         Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	00 mg
Collectionaria			
(2-Hydroxybenzoic acid)	Cat. No.: HY-B0167	(2-Hydroxybenzoic acid-d6)	Cat. No.: HY-B0167S
Salicylic acid (2-Hydroxybenzoic acid) inhibits cyclo-oxygenase-2 ( <b>COX-2</b> ) activity independently of transcription factor (NF-κB) activation.	ОН	Salicylic acid-D6 (2-Hydroxybenzoic acid-D6) is a deuterium labeled Salicylic acid. Salicylic acid inhibits cyclo-oxygenase-2 ( <b>COX-2</b> ) activity independently of transcription factor (NF-κB) activation.	
Purity:         96.22%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 500 mg, 10 g, 50 g	√ `он	Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg	Ď
Salidroside (Rhodioloside)	<b>Cat. No.:</b> HY-N0109	Salinomycin (Procoxacin)	<b>Cat. No.:</b> HY-15597
Salidroside is a <b>prolyl endopeptidase</b> inhibitor. Salidroside alleviates cachexia symptoms in mouse models of cancer cachexia via activating <b>mTOR</b> signalling. Salidroside protects dopaminergic neurons by enhancing PINK1/Parkin-mediated mitophagy.	но сторон сон	Salinomycin (Procoxacin), a polyether potassium ionophore antibiotic, selectively inhibits the growth of gram-positive bacteria. Salinomycin is a potent inhibitor of Wnt/ $\beta$ -catenin signaling, blocks Wnt-induced LRP6 phosphorylation.	
Purity:99.79%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg		Purity:     ≥98.0%       Clinical Data:     No Development Reported       Size:     10 mM × 1 mL, 5 mg, 10 mg, 50 mg	

Salinomycin sodium salt		Salubrinal
(Salinomycin sodium; Sodium salinomycin)	Cat. No.: HY-17439	<b>Cat. No.:</b> HY-15486
Salinomycin sodium salt (Salinomycin sodium), an antibiotic potassium ionophore, is a potent inhibitor of <b>Wnt/β-catenin</b> signaling.		Salubrinal is a cell-permeable and selective inhibitor of eIF2 $\alpha$ dephosphorylation. Salubrinal acts as a dual-specificity phosphatase 2 (Dusp2) inhibitor and suppresses inflammation in anti-collagen antibody-induced arthritis.
Purity: >98% Clinical Data: No Development Reported		Purity: 99.69% Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 25 mg, 50 mg, 100 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg
Samuraciclib		Samuraciclib hydrochloride
(CT7001; ICEC0942)	Cat. No.: HY-103712	(CT7001 hydrochloride; ICEC0942 hydrochloride) Cat. No.: HY-103712A
Samuraciclib (CT7001) is a potent, selective, ATP-competitive and orally active <b>CDK7</b> inhibitor, with an $IC_{50}$ of 41 nM. Samuraciclib displays 45-, 15-, 230- and 30-fold selectivity over CDK1, CDK2 (IC ₅₀ of 578 nM), CDK5 and CDK9, respectively.	NH NH N N	Samuraciclib hydrochloride (CT7001 hydrochloride) is a potent, selective, ATP-competitive and orally active CDK7 inhibitor, with an IC ₅₀ of 41 nM. Samuraciclib hydrochloride displays 45-, 15-, 230- and 30-fold selectivity over CDK1, CDK2 (IC ₅₀ of 578 nM), CDK5 and CDK9, respectively.
Purity: >98%		Purity: 99.98% H-Cl
Clinical Data: No Development Reported		Clinical Data: Phase 2 Size: 10 mM × 1 mL 5 mg 10 mg 25 mg 50 mg 100 mg
512C. 1 mg, 5 mg		Size. 10 mill + 1 mil, 5 mg, 20 mg, 20 mg, 300 mg
Samuraciclib hydrochloride hydrate (CT7001 hydr	ochloride	Sandacanol
hydrate; ICEC0942 hydrochloride hydrate)	Cat. No.: HY-103712B	Cat. No.: HY-N7707
Samuraciclib (CT7001) hydrochloride hydrate is a potent, selective, ATP-competitive and orally active <b>CDK7</b> inhibitor, with an <b>IC</b> ₅₀ of 41 nM.		Sandacanol is a specific agonist of <b>olfactory</b> <b>receptor</b> ( <b>OR10H1</b> ). Sandacanol induces cell cycle arrest and some <b>apoptosis</b> in bladder cancer cells. HO
Purity: 99.08% Clinical Data: Phase 2	2.5 HCI H2O H2O	Purity: >98% Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	.00 mg	Size: 100 mg
Sanguinarine		Sanguinarine chloride (Sanguinarin chloride; Sanguinarium
(Sanguinarin; Sanguinarium; Pseudochelerythrine)	Cat. No.: HY-N0052	chloride; Pseudochelerythrine chloride) Cat. No.: HY-N0052A
Sanguinarine (Sanguinarin), a benzophenanthridine alkaloid derived from the root of Sanguinaria Canadensis, can stimulate <b>apoptosis</b> via activating the production of reactive oxygen species (ROS). Sanguinarine-induced apoptosis is associated with the activation of JNK and NF-κB. <b>Purity:</b> >98% <b>Clinical Data:</b> No Development Reported <b>Size:</b> 5 mg, 10 mg, 25 mg	STT - So	Sanguinarine (Sanguinarin) chloride, a benzophenanthridine alkaloid derived from the root of Sanguinaria Canadensis, can stimulate <b>apoptosis</b> via activating the production of reactive oxygen species (ROS). Sanguinarine-induced apoptosis is associated with the activation of JNK and NF- $\kappa$ B. <b>Purity:</b> 99.24% <b>Clinical Data:</b> No Development Reported <b>Size:</b> 10 mM × 1 mL, 5 mg, 10 mg
Connenshelson		CAD105044
Sappanchaicone	Cat No : HV-59001	SAK123644
Sappanchalcone, a flavonoid isolated from Caesalpinia sappan L., induces caspase-dependent and AIF-dependent <b>apoptosis</b> in human colon cancer cells.		SAR125844 is a potent, highly selective, reversible and ATP-competitive <b>MET receptor</b> <b>tyrosine kinase (RTK)</b> inhibitor, with an IC ₅₀ of 4.2 nM. Shows inhibition of MET autophosphorylation in cell-based assays.
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg		Purity:98.11%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg



SC66		SC99	
	Cat. No.: HY-19832		Cat. No.: HY-124858
SC66 is an <b>Akt</b> inhibitor, reduces cell viability in a dose- and time-dependent manner, inhibits colony formation and induces apoptosis in hepatocellular carcinoma (HCC) cells.	NT CN	SC99 is an orally active, selective <b>STAT3</b> inhibitor targeting JAK2-STAT3 pathway. SC99 docks into the ATP-binding pocket of JAK2. SC99 inhibits phosphorylation of JAK2 and STAT3 with no effects on the other kinases associated with STAT3 signaling.	
Purity:99.88%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg		Purity:         99.07%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
SCH79797	<b>Cat. No.:</b> HY-14993	SCH79797 dihydrochloride	<b>Cat. No.:</b> HY-14994
SCH79797 is a highly potent, selective nonpeptide protease activated receptor 1 (PAR1) antagonist. SCH79797 inhibits binding of a high-affinity thrombin receptor-activating peptide to PAR1 with an $IC_{s0}$ of 70 nM and a $K_i$ of 35 nM.		SCH79797 dihydrochloride is a highly potent, selective nonpeptide <b>protease activated receptor</b> <b>1 (PAR1)</b> antagonist. SCH79797 dihydrochloride inhibits binding of a high-affinity thrombin receptor-activating peptide to <b>PAR1</b> with an $IC_{50}$ of 70 nM and a K ₁ of 35 nM.	
Clinical Data:       No Development Reported         Size:       5 mg, 10 mg, 50 mg, 100 mg		Clinical Data:       No Development Reported         Size:       1 mg, 5 mg, 10 mg	
Schinifoling		Schisandrin C	
Schimoline	Cat. No.: HY-N4164	(Schizandrin-C; Wuweizisu-C)	<b>Cat. No.:</b> HY-N0690
Schinifoline, a 4-quinolinone derivative isolated from Zanthoxylum schinifolium Sieb, improves radiosensitizing effect, and effects cell cycle and apoptotic-inducing effects in cancer.	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	Schisandrin C (Schizandrin-C) is a phytochemical <b>lignan</b> isolated from Schizandra chinensis. Schisandrin C has diverse biological activities, including anticancer, anti-inflammatory and antioxidant effects.	
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:99.95%Clinical Data:No Development ReportedSize:10 mg, 50 mg, 100 mg	0 ~ ~
Sclareol	<b>Cat. No.:</b> HY-N0128	Scopoletin (Gelseminic acid; Chrysatropic acid)	<b>Cat. No.:</b> HY-N0342
Sclareol is isolated from Salvia sclarea with anticarcinogenic activity. Sclareol shows strong cytotoxic activity against mouse leukemia (P-388), human epidermal carcinoma (KB) cells and human leukemia cell lines. Purity: ≥98.0% Clinical Data: No Development Reported	OH OH	Scopoletin is an inhibitor of acetylcholinesterase (AChE). Purity: 99.70% Clinical Data: No Development Reported	HO
Size: 25 mg, 50 mg, 100 mg		Size: 50 mg, 100 mg, 200 mg	
Scoulerine ((-)-Scoulerine; Discretamine)	<b>Cat. No.:</b> HY-N1255	SCR130	<b>Cat. No.</b> : HY-139297
Scoulerine ((-)-Scoulerine), an isoquinoline alkaloid, is a potent <b>antimitotic</b> compound. Scoulerine is also an inhibitor of <b>BACE1</b> (β-site amyloid precursor protein cleaving enzyme 1). Scoulerine inhibits proliferation, arrests cell cycle, and induces apoptosis in cancer cells.	OH OH OH	SCR130 is a SCR7-based <b>DNA nonhomologous</b> end-joining (NHEJ) inhibitor. SCR130 inhibits the end-joining of DNA in a Ligase IV-dependent manner. SCR130 is specific to Ligase IV, and shows minimal or no effect on Ligase III and Ligase I mediated joining.	
Purity:     99.27%       Clinical Data:     No Development Reported       Size:     1 mg		Purity:         98.00%           Clinical Data:         No Development Reported           Size:         5 mg, 10 mg	CI

SCR7	<b>Cat. No.:</b> HY-12742	SCR7 pyrazine	<b>Cat. No.:</b> HY-107845
SCR7 is an unstable form that can be autocyclized into a stable form SCR7 pyrazine. SCR7 pyrazine is a DNA ligase IV inhibitor that blocks nonhomologous end-joining (NHEJ) in a ligase IV-dependent manner.         Purity:       98.22%         Clinical Data:       No Development Reported         Size:       5 mg		SCR7 pyrazine is a DNA ligase IV inhibitor that blocks nonhomologous end-joining (NHEJ) in a ligase IV-dependent manner. SCR7 pyrazine is also a CRISPR/Cas9 enhancer which increases the efficiency of Cas9-mediated homology-directed repair (HDR).         Purity:       98.70%         Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg	
Scriptaid		SD-1008	
(Scriptide; GCK1026)	Cat. No.: HY-15489		Cat. No.: HY-107595
Scriptaid is a potent histone deacetylase (HDAC) inhibitor, used in cancer research. Scriptaid is also a sensitizer to antivirals and has potential for epstein-barr virus (EBV)-associated lymphomas treatment. Purity: 98.59% Clinical Data: No Development Reported Size: 10 mM × 1 mL 10 mg. 50 mg	C HN-OH	SD-1008 is a potent JAK inhibitor. SD-1008 inhibits tyrosyl phosphorylation of STAT3, JAK2 and Src. SD-1008 also reduces STAT3-dependent luciferase activity. SD-1008 enhances <b>apoptosis</b> induced by Paclitaxel in ovarian cancer cells via directly blocking the JAK-STAT3 signaling pathway. Purity: >98% Clinical Data: No Development Reported Size: 1 mg. 5 mg	
		5. 5	
SD-36	<b>Cat. No.:</b> HY-129602	Se-Methylselenocysteine (Methylselenocysteine; Se-Methylseleno-L-cysteine)	<b>Cat. No.</b> : HY-114245
SD-36 is a potent and efficacious STAT3 PROTAC degrader ( $K_d = \sim 50$ nM), and demonstrates high selectivity over other STAT members. SD-36 also effectively degrades mutated STAT3 proteins in cells and suppresses the transcriptional activity of STAT3 ( $IC_{so}=10$ nM). Purity: 99.46% Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg		Se-Methylselenocysteine, a precursor of Methylselenol, has potent cancer chemopreventive activity and anti-oxidant activity.         Se-Methylselenocysteine is orally bioavailable, and induces <b>apoptosis</b> .         Purity:       ≥98.0%         Clinical Data:       Phase 1         Size:       10 mM × 1 mL, 5 mg, 10 mg	Se NH2
SEC	<b>Cat. No.:</b> HY-125355	SecinH3	<b>Cat. No</b> .: HY-100559
SEC induces activation of <b>ANXA7 GTPase</b> via the AMPK/mTORC1/STAT3 signaling pathway. SEC selectively promotes <b>apoptosis</b> in cancer cells, expressing a high level of ITGB4 by inducing ITGB4 nuclear translocation.	- C N OH	SecinH3 is an antagonist of <b>cytohesins</b> with $IC_{s0}$ s of 5.4 $\mu$ M, 2.4 $\mu$ M, 5.4 $\mu$ M, 5.6 $\mu$ M, 5.6 $\mu$ M and 65 $\mu$ M for hCyh1, hCyh2, mCyh3, hCyh3, drosophila steppke and yGea2-S7, respectively.	Provide Contraction
Purity:         98.13%           Clinical Data:         Phase 1           Size:         10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	10 mg	Purity:99.54%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg	0
SEL24-B489	<b>Cat. No.</b> : HY-120758	Selonsertib (GS-4997)	<b>Cat. No.:</b> HY-18938
SEL24-B489 is a potent, type I, orally active, dual <b>PIM</b> and <b>FLT3-ITD</b> inhibitor, with $K_d$ values of 2 nM for PIM1, 2 nM for PIM2 and 3 nM for PIM3, respectively. .		Selonsertib (GS-4997), an orally bioavailable, selective apoptosis signal-regulating kinase 1 (ASK1) inhibitor with a $pIC_{s0}$ of 8.3, has been evaluated as an experimental treatment for diabetic nephropathy and kidney fibrosis.	
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	0.0	Purity:         98.99%           Clinical Data:         Phase 2           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 70	200 mg

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SHR0302	Cot No. UV 112724	SID 3712249	Cot No. 11V 10721
SHR0302 is a potent and orally active all members of the JAK family inhibitor, particularly JAK1. The selectivity of SHR0302 for JAK1 is >10-fold for JAK2, 77-fold for JAK3, 420-fold for Tyk2. Purity: 99.58% Clinical Data: No Development Reported		SID 3712249 (MiR-544 Inhibitor 1) is an inhibitor of the biogenesis of microRNA-544 (miR-544). Target: MiR-544 MiR-544 represses expression of mTOR, promoting tumor cell survival in a hypoxic environment. Purity: 98.35% Clinical Data: No Davidocement Reported	
Size: 5 mg, 10 mg, 25 mg, 50 mg		Size:         10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 3	LOO mg
Sideroxylin	<b>Cat. No.:</b> HY-N1306	Sildenafil (UK-92480)	<b>Cat. No.:</b> HY-15025
Sideroxylin is a C-methylated flavone isolated from Callistemon lanceolatus and exerts antimicrobial activity against <b>Staphylococcus aureus</b> .	OH OF OH	Sildenafil (UK-92480) is a potent phosphodiesterase type 5 (PDE5) inhibitor with an $IC_{50}$ of 5.22 nM.	N N O HN N N N O HN N O V O V
Purity:     >98%       Clinical Data:     No Development Reported       Size:     5 mg		Purity:         99.90%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 50 mg, 100 mg, 200 mg	
Sildenafil citrate		Sildenafil-d8	
(UK-92480 citrate)	Cat. No.: HY-15025A	(UK-92480-d8)	Cat. No.: HY-15025S1
Sildenafil citrate is a potent phosphodiesterase type 5 ( <b>PDE5</b> ) inhibitor with $IC_{so}$ of 5.22 nM.	N N. O HN IN O HN IN	Sildenafil-d8 (UK-92480-d8) is the deuterium labeled Sildenafil. Sildenafil (UK-92480) is a potent phosphodiesterase type 5 ( <b>PDE5</b> ) inhibitor with an $IC_{50}$ of 5.22 nM.	N P P N N N N N N N N N N N N N N N N N
Purity:         99.73%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 50 mg, 100 mg, 200 mg, 500 r	ноцонон	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 10 mg	~~~ J
Silvestrol		Silybin	
((-)-Silvestrol)	Cat. No.: HY-13251		Cat. No.: HY-N0779A
Silvestrol is a eukaryotic translation initiation factor 4A ( <b>eIF4A</b> ) inhibitor isolated from the fruits and twigs of Aglaia foveolata. Silvestrol induces <b>autophagy</b> and caspase-mediated <b>apoptosis</b> .	но Эсторо Сорона С	Silybin is a flavonolignan isolated from milk thistle (Silybum marianum) seeds. Silybin induces <b>apoptosis</b> and exhibits hepatoprotective, antioxidant, anti-inflammatory, anti-cancer activity.	но стор он он он
Purity:98.11%Clinical Data:No Development ReportedSize:1 mg, 2 mg, 5 mg, 10 mg		Purity:>98%Clinical Data:Phase 4Size:5 mg, 10 mg, 25 mg	
Simvastatin		Simvastatin-d6	
(MK 733)	Cat. No.: HY-17502	(MK 733-d6)	Cat. No.: HY-110231
Simvastatin (MK 733) is a competitive inhibitor of HMG-CoA reductase with a $K_i$ of 0.2 nM.		Simvastatin-d6 (MK 733-d6) is the deuterium labeled Simvastatin. Simvastatin (MK 733) is a competitive inhibitor of <b>HMG-CoA reductase</b> with a $K_i$ of 0.2 nM.	
Purity:         99.45%           Clinical Data:         Launched           Size:         50 mg, 100 mg, 200 mg, 500 mg	and the second sec	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	* * *
Sinapinic acid		Sinomenine	
--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------	------------------------------	-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------	-----------------------------------------------------------------
(Sinapic acid)	Cat. No.: HY-W009732		Cat. No.: HY-15122
Sinapinic acid (Sinapic acid) is a phenolic compound isolated from Hydnophytum formicarum Jack. Rhizome, acts as an inhibitor of HDAC, with an $IC_{so}$ of 2.27 mM, and also inhibits ACE-I activity.Purity:99.77%	от страна но страна он	Sinomenine, an alkaloid extracted from Sinomenium acutum, is a blocker of the NF-κB activation. Sinomenine also is an activator of μ-opioid receptor. Purity: 99.88%	OCH OC
Clinical Data:No Development ReportedSize:10 mM × 1 mL, 100 mg		Clinical Data: Launched Size: 10 mM × 1 mL, 100 mg	
Sinomenine hydrochloride (Cucoline hydrochloride)	Cat. No.: HY-15122A	Siomycin A	Cat. No.: HY-P1687
Sinomenine hydrochloride (Cucoline hydrochloride), an alkaloid extracted from Sinomenium acutum, is a blocker of the NF- $\kappa$ B activation. Sinomenine also is an activator of $\mu$ -opioid receptor.	OCH OC	Siomycin A is a thiopeptide antibiotic and is a Forkhead box M1(FOXM1) selective inhibitor without affecting other members of the Forkhead box family. Siomycin A has anti-tumor and promotes apoptosis.	
Purity:         99.88%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 100 mg	HCI	Purity:     >98%       Clinical Data:     No Development Reported       Size:     500 μg	τ _μ τ.
SIRT7 inhibitor 97491		Sirtinol	
	Cat. No.: HY-135899		Cat. No.: HY-13515
SIRT7 inhibitor 97491, a potent <b>SIRT7</b> inhibitor with an IC ₅₀ of 325 nM, reduces deacetylase activity of SIRT7 in a dose-dependent manner. SIRT7 inhibitor 97491 prevents tumor progression by increasing p53 stability through acetylation at K373/382.		Sirtinol is a <b>sirtuin (SIRT)</b> inhibitor, with $IC_{so}s$ of 48 $\mu$ M, 57.7 $\mu$ M and 131 $\mu$ M for ySir2, hSIRT2 and hSIRT2, respectively.	Ç, t , c , f oH
Purity:         98.05%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity:     ≥98.0%       Clinical Data:     No Development Reported       Size:     10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	~ ~
SKF-96365 hydrochloride		SKI II	
	Cat. No.: HY-100001		Cat. No.: HY-13822
SKF-96365 hydrochloride is a potent <b>TRP channel</b> blocker and a <b>store-operated Ca²⁺ entry (SOCE)</b> inhibitor. SKF-96365 hydrochloride significantly inhibits hERG, hKCNQ1/hKCNE1, hKir2.1 and hKv4.3 current, and significantly prolongs the QTc interval in isolated guinea pig hearts. <b>Purity:</b> 99.51% <b>Clinical Data:</b> No Development Reported		SKI-II is an oral active and synthetic inhibitor of sphingosine kinase (SK) activity, with IC _{so} values of 78 μM and 45 μM for SK1 and for SK2, respectively. SKI II causes an irreversible inhibition of SK1 by inducing its lysosomal and/or proteasomal degradation.         Purity:       99.88%         Clinical Data:       No Development Reported	CI C
Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 2	200 mg	Size: 10 mM × 1 mL, 10 mg, 25 mg, 50 mg, 100 mg,	200 mg
SKI V	<b>Cat. No</b> .: HY-12895	SKI-178	<b>Cat. No.:</b> HY-12892
SKI V is a noncompetitive and potent non-lipid sphingosine kinase (SPHK; SK) inhibitor with an $IC_{so}$ of 2 $\mu$ M for GST-hSK. SKI V potently inhibits PI3K with an $IC_{so}$ of 6 $\mu$ M for hPI3k. SKI V decreases formation of the mitogenic second messenger sphingosine-1-phosphate (S1P). Purity: 98.09% Clinical Data: No Development Reported	ССС-ОН ОН	SKI-178 is a potent sphingosine kinase-1 (SphK1) and SphK2 inhibitor. SKI-178 is cytotoxic at IC ₅₀ concentrations ranging from 1.8 to 0.1 µM in both drug sensitive and multi-drug resistant cancer cell lines (i.e., MTR3, NCI-ADR and HL60/VCR cells).         Purity:       98.05%         Clinical Data:       No Development Reported	~()- ₆ + ¹ # ^N + ⁽⁾
Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	0 mg

SKI-I	C + N + 10/ 115725	SM-164	C + N - 11/ 15000
SKI-I is a potent and selective inhibitor of human sphingosine kinase (SK), with an IC ₅₀ of 1.2 $\mu$ M for ST-hSK. SKI-I also inhibits hERK2 (IC ₅₀ =11 $\mu$ M). SKI-I induces apoptosis in tumor cell lines.	Cat. No.: HY-115735	SM-164 is a cell-permeable Smac mimetic compound. SM-164 binds to XIAP protein containing both the BIR2 and BIR3 domains with an $IC_{so}$ value of 1.39 nM and functions as an extremely potent antagonist of XIAP.	Cat. No.: HY-15989
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 50 mg, 100 mg		Purity:99.65%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 50 mg, 100 mg	
SM-164 Hydrochloride	Cat No : HV-15989A	SMIP004	Cat No: HV-15694
SM-164 Hydrochloride is a cell-permeable Smac mimetic compound. SM-164 binds to XIAP protein containing both the BIR2 and BIR3 domains with an $IC_{s0}$ value of 1.39 nM and functions as an extremely potent antagonist of XIAP.	+ 20	SMIP004 is a <b>SKP2 E3 ligase</b> inhibitor, which downregulates SKP2 and to stabilise p27. SMIP004 is a cancer cell selective <b>apoptosis</b> inducer of human prostate cancer cells.	
Purity:99.0%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg		Purity:98.66%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 50 mg, 100	) mg
SNS-032 (BMS-387032)	Cat No : HV-10008	Sodium 4-phenylbutyrate (4-PBA sodium; 4-Phenylbu	utyric acid
SNS-032 (BMS-387032) is a potent and selective inhibitor of CDK2, CDK7, and CDK9 with $IC_{50}$ s of 38 nM, 62 nM and 4 nM, respectively. SNS-032 has antitumor effect.		Sodium 4-phenylbutyrate (4-PBA sodium) is an inhibitor of HDAC and endoplasmic reticulum (ER) stress, used in cancer and infection research.	
Purity:         99.49%           Clinical Data:         Phase 1           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 2	200 mg	Purity:99.96%Clinical Data:LaunchedSize:100 mg, 200 mg	
Odium diatrizoate (Diatrizoic acid sodium salt; Sodium amidotrizoate)	Cat. No.: HY-B0926A	Sodium dichloroacetate	<b>Cat. No.</b> : HY-Y0445A
Sodium diatrizoate (Diatrizoic acid sodium salt) is an iodinated radiocontrast agent and has the potential for radiographic imaging of the airways. Sodium diatrizoate induces mitochondrial turnover and oxidative stress, and activating apoptosis by dysregulating calcium. Purity: ≥98.0% Clinical Data: No Development Reported		Sodium dichloroacetate is a metabolic regulator in cancer cells' mitochondria with anticancer activity. Sodium dichloroacetate inhibits PDHK, resulting in decreased lactic acid in the tumor microenvironment. Purity: ≥98.0% Clinical Data: Phase 3 Size: 100 mg	CI CI CI
512C. 10 IIIVI A 1 IIIL, 300 IIIY		5126. 100 mg	
Sodium oleate (Oleic acid sodium; 9-cis-Octadecenoic a sodium; 9Z-Octadecenoic acid sodium)	acid Cat. No.: HY-N1446B	Sodium Salicylate (Salicylic acid sodium salt; 2-Hydroxybenzoic acid sodium salt)	<b>Cat. No.:</b> HY-B0167A
Sodium oleate (Oleic acid sodium) is an abundant monounsaturated fatty acid sodium. Sodium oleate is a Na ⁺ /K ⁺ ATPase activator.	low	Sodium Salicylate (Salicylic acid sodium salt) inhibits cyclo-oxygenase-2 (COX-2) activity independently of transcription factor (NF-ĸB) activation. Sodium Salicylate is also a S6K inhibitor.	ONa
Purity:     ≥98.0%       Clinical Data:     Launched       Size:     10 mM × 1 mL, 100 mg		Purity:         99.88%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 500 mg, 10 g, 50 g	~ ОН

Solamargine		Solasodine	
(Solamargin; δ-Solanigrine)	Cat. No.: HY-N0069	(Purapuridine; Solancarpidine; Solasodin)	Cat. No.: HY-N0068
Solamargine, a derivative from the steroidal solasodine in Solanum species, exhibits anticancer activities in numerous types of cancer. Solamargine induces non-selective cytotoxicity and <b>P-glycoprotein</b> inhibition. <b>Purity:</b> ≥98.0% <b>Clinical Data:</b> No Development Reported	st Starte	Solasodine (Purapuridine) is a steroidal alkaloid that occurs in plants of the Solanaceae family. Solasodine has neuroprotective, antifungal, hypotensive, anticancer, antiatherosclerotic, antiandrogenic and anti-inflammatory activities. Purity: 98.86% Clinical Data: No Development Reported	HO HH H
Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg		Size: 10 mg, 50 mg, 100 mg	
Sophocarpine	<b>Cat. No.</b> : HY-N0103	Sophocarpine monohydrate	<b>Cat. No.:</b> HY-N0103A
Sophocarpine is one of the significant alkaloid extracted from the traditional herb medicine Sophora flavescens which has many pharmacological properties such as anti-virus, anti-tumor, anti-inflammatory.		Sophocarpine (monohydrate) is one of the significant alkaloid extracted from the traditional herb medicine Sophora flavescens which has many pharmacological properties such as anti-virus, anti-tumor, anti-inflammatory.	
Purity:     ≥98.0%       Clinical Data:     No Development Reported       Size:     10 mM × 1 mL, 5 mg, 10 mg, 20 mg	0	Purity:       99.15%         Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 5 mg, 10 mg	H ₂ O
Sophoraflavanone G		Sophoridine	
(Kushenol F)	Cat. No.: HY-N1231		Cat. No.: HY-N1373
Sophoraflavanone G (Kushenol F) is iaolated from Sophora flavescens and shows anti-tumor and anti-inflammatory properties. Sophoraflavanone G (Kushenol F) induces MDA-MB-231 and HL-60 cells <b>apoptosis</b> through suppression of MAPK-related pathways. <b>Purity:</b> 98.30%	но со он	Sophoridine is a quinolizidine alkaloid isolated from leafs of Leguminous plant Sophora alopecuroides.L. Sophoridine induces <b>apoptosis</b> . Sophoridine has the potential to be a novel, potent and selective antitumor drug candidate for pancreatic cancer with well-tolerated toxicity. Purity: ≥98.0%	
Size: 5 mg, 10 mg, 20 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 20 mg	
Sorafenib		Sorafenib Tosylate	
Sorafenib (Bay 43-9006) is a potent and orally active <b>Raf</b> inhibitor with $IC_{so}$ of 6 nM and 20 nM for <b>Raf-1</b> and <b>B-Raf</b> , respectively. Sorafenib is a multikinase inhibitor with $IC_{so}$ of 90 nM, 15 pM 20 pM 57 nM and 58 nM for VEGED VEGED		Sorafenib Tosylate (Bay 43-9006 Tosylate) is a potent and orally active <b>Raf</b> inhibitor with IC ₅₀ s of 6 nM and 20 nM for <b>Raf-1</b> and <b>B-Raf</b> , respectively.	Sayora'r A
PDGFRβ, FLT3 and c-Kit, respectively.         Purity:       99.92%         Clinical Data:       Launched         Size:       10 mM × 1 mL, 100 mg, 500 mg		Purity:         99.75%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 100 mg, 500 mg	
Sorafenib-13C,d3		Sorafenib-d3	
·	Cat. No.: HY-10201S2	(Bay 43-9006-d3; Donafenib)	Cat. No.: HY-10201S
Sorafenib-13C,d3 is the 13C- and deuterium labeled Sorafenib. Sorafenib (Bay 43-9006) is a potent and orally active Raf inhibitor with $IC_{so}$ of 6 nM and 20 nM for Raf-1 and B-Raf, respectively.		Sorafenib-d3 (Bay 43-9006-d3) is the deuterium labeled Sorafenib. Sorafenib is a multikinase inhibitor $IC_{s0}$ s of 6 nM, 20 nM, and 22 nM for Raf-1, B-Raf, and VEGFR-3, respectively.	pri corte
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:         99.57%           Clinical Data:         Launched           Size:         5 mg, 10 mg, 25 mg, 50 mg, 100 mg	

Sorafenib-d4		Soyasapogenol A	
(Bay 43-9006-d4)	Cat. No.: HY-10201S1		Cat. No.: HY-N6073
Sorafenib-d4 (Bay 43-9006-d4) is the deuterium labeled Sorafenib. Sorafenib is a multikinase inhibitor IC ₅₀ s of 6 nM, 20 nM, and 22 nM for Raf-1, B-Raf, and VEGFR-3, respectively.	2049. grade	Soyasapogenol A, a triterpene compound, isolated from the roots of Abrus cantoniensis.	но 1
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:99.06%Clinical Data:No Development ReportedSize:5 mg	он
Soyasapogenol B	<b>Cat. No.:</b> HY-N6074	Soyasaponin III	<b>Cat. No.:</b> HY-N7273
Soyasapogenol B, an ingredient of soybean, exerts anti-proliferative, anti-metastatic activities. Soyasapogenol B triggers endoplasmic reticulum stress, which mediates <b>apoptosis</b> and <b>autophagy</b> in colorectal cancer.		Soyasaponin III, a monodesmodic oleanane triterpenoid, is one of the main potentially bioactive saponins found in soy (Glycine max) and related products. Soyasaponin III can induce <b>apoptosis</b> in Hep-G2 cells. <b>Purity:</b> 97.72%	
Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg		Clinical Data: No Development Reported Size: 1 mg	Če+
SP2509		SP600125	
51 2505	Cat. No.: HY-12635	51 000125	Cat. No.: HY-12041
SP2509 is a potent and selective antagonist of lysine specific demethylase 1 (LSD1) with an IC ₅₀ of 13 nM.	a Constant of the second	SP600125 is an orally active, reversible, and ATP-competitive JNK inhibitor with IC ₅₀ s of 40, 40 and 90 nM for JNK1, JNK2 and JNK3, respectively. SP600125 is a potent <b>ferroptosis</b> inhibitor. SP600125 inhibits <b>autophagy</b> and activates apartments	N-NH
Purity:99.90%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity:       99.55%         Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg/time	g, 500 mg
Sparfosic acid	<b>Cat. No.</b> : HY-112732	Sparfosic acid trisodium	<b>Cat. No.:</b> HY-112732B
Sparfosic acid, a DNA antimetabolite agent, is a potent inhibitor of aspartate transcarbamoyl transferase, the enzyme catalyzing the second step of de novo pyrimidine biosynthesis.	о о о о он но он н он	Sparfosic acid trisodium is a DNA antimetabolite agent and a potent inhibitor of <b>aspartate</b> <b>transcarbamoyl transferase</b> . Aspartate transcarbamoyl transferase catalyzes the second step of de novo pyrimidine biosynthesis.	
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 50 mg, 100 mg		Purity:         99.65%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
Spautin-1	<b>Cat. No.:</b> HY-12990	SR-4835	<b>Cat. No.:</b> HY-130250
Spautin-1 is a specific and potent <b>autophagy</b> inhibitor which inhibits ubiquitin-specific peptidases, USP10 and USP13 with <b>IC</b> ₅₀ s of 0.6-0.7 μM.	F NH F	SR-4835 is a potent, highly selective and ATP competitive dual inhibitor of <b>CDK12/CDK13</b> (CDK12: $IC_{s0}$ =99 nM, $K_a$ =98 nM; CDK13: $K_a$ =4.9 nM). SR-4835 acts in synergy with DNA-damaging chemotherapy and PARP inhibitors and provokes triple-negative breast cancer (TNBC) cell death.	
Purity:         99.26%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity:         99.82%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	_ک—< 00 mg

## SRT 2183 **SS28** Cat. No.: HY-19759 Cat. No.: HY-100761 SRT 2183 is a selective Sirtuin-1 (SIRT1) activator SS28, a SRT501 analog with oral bioavailability, with an $\text{EC}_{1.5}$ value of 0.36 $\mu\text{M}.$ SRT 2183 induces inhibits tubulin polymerization to cause cell growth arrest and apoptosis, concomitant with cycle arrest at G2/M phase. SS28 results in deacetylation of STAT3 and NF-KB, and reduction of apoptosis rather than necrosis tubulin. c-Myc protein levels. 98.48% >98% Purity: Purity: Clinical Data: No Development Reported Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg Size: 1 mg, 5 mg SSE15206 SSK1 Cat. No.: HY-111425 Cat. No.: HY-138936 SSE15206 is a microtubule polymerization SSK1, a senescence-specific killing compound, is a inhibitor (GI₅₀ = 197 nM in HCT116 cells) that $\beta$ -galactosidase-targeted prodrug attenuates overcomes multidrug resistance. Causes aberrant inflammation. SSK1 is activated by lysosomal mitosis resulting in G2/M arrest due to incomplete β-galactosidase and selectively killed senescent spindle formation in cancer cells. cells through the activation of p38 MAPK and induction of apoptosis. Purity: 98 39% **Purity:** 99 1 9% Clinical Data: No Development Reported Clinical Data: No Development Reported 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg Size: Size: 5 mg, 10 mg STAT3-IN-1 Stachyose tetrahydrate Cat. No.: HY-113529 Cat. No.: HY-100753 STAT3-IN-1 (compound 7d) is an excellent, Stachyose tetrahydrate, a functional oligosaccharide, acts as a prebiotic. Stachyose selective and orally active STAT3 inhibitor, with tetrahydrate can prevent indirectly colon cancer $\text{IC}_{\text{so}}$ values of 1.82 $\mu\text{M}$ and 2.14 $\mu\text{M}$ in HT29 and MDA-MB 231 cells, respectively. STAT3-IN-1 cell growth by promoting the proliferation of probiotics or producing beneficial materials in (compound 7d) induces tumor apoptosis. the intestine. Purity: 98.10% 96.54% Purity: Clinical Data: No Development Reported Clinical Data: No Development Reported 5 mg, 10 mg, 25 mg Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg Size: STAT3-IN-10 STAT3-IN-3 Cat. No.: HY-146728 Cat. No.: HY-128588 STAT3-IN-10 (A11) is a STAT3 inhibitor with an STAT3-IN-3 is a potent and selective inhibitor of IC_{so} value of 5.18 µM. STAT3-IN-10 directly binds signal transducer and activator of transcription 3 to STAT3 SH2 domain, inhibits tumor cell growth (STAT3), with anti-proliferative activity. and induces apoptosis in cancer cells. STAT3-IN-3 induces apoptosis in breast cancer cells. Purity: **Purity:** 98.23% >98% Clinical Data: No Development Reported Clinical Data: No Development Reported Size: 1 mg, 5 mg Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg STAT5-IN-2 Stattic Cat. No.: HY-102048 Cat. No.: HY-13818 STAT5-IN-2 is a STAT5 inhibitor, extracted from Stattic is a potent STAT3 inhibitor and inhibits reference 1, example 17f. STAT5-IN-2 has potent STAT3 phosphorylation (at Y705 and S727). Stattic antileukemic effect. inhibits the binding of a high affinity phosphopeptide for the SH2 domain of STAT3. Stattic ameliorates the renal dysfunction in Alport syndrome (AS) mice. Purity: 99.01% Purity: ≥97.0% No Development Reported Clinical Data: No Development Reported Clinical Data: Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg 10 mM × 1 mL, 10 mg, 25 mg, 50 mg, 100 mg Size:



Suberoyl bis-hydroxamic acid	Cat No : HY-W009776	Sulfasalazine	Cat No: HY-14655
Suberoyl bis-hydroxamic acid (Suberohydroxamic acid; SBHA) is a competitive and cell-permeable HDAC1 and HDAC3 inhibitor with ID ₅₀ values of 0.25 µM and 0.30 µM, respectively.	но Ц он ц он	Sulfasalazine (NSC 667219) is an anti-rheumatic agent for the research of rheumatoid arthritis and ulcerative colitis. Sulfasalazine can suppress NF-κB activity. Sulfasalazine is a type 1 ferroptosis inducer.	
Purity:     ≥98.0%       Clinical Data:     No Development Reported       Size:     10 mM × 1 mL, 50 mg, 100 mg, 250 mg		Purity:         99.04%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 500 mg, 1 g, 5 g	
Sulfasalazine-d4	<b>Cat. No.</b> : HY-14655S	Sulforaphane	<b>Cat. No.:</b> HY-13755
Sulfasalazine-d4 is the deuterium labeled Sulfasalazine. Sulfasalazine (NSC 667219) is an anti-rheumatic agent for the research of rheumatoid arthritis and ulcerative colitis. Sulfasalazine can suppress NF-kB activity. Sulfasalazine is a type 1 ferroptosis inducer. Purity: >98% Clinical Data: No Development Reported	of the state of th	Sulforaphane is an isothiocyanate present naturally in widely consumed vegetables. Sulforaphane increases tumor suppressor protein transcription and inhibits <b>histone deacetylase</b> activity. <b>Purity:</b> 99.75% Clinical Data: Phase 3	s _{sccsN}
Size: 2.5 mg, 25 mg		Size:         10 mg, 25 mg, 50 mg, 100 mg	
Sulforaphene	<b>Cat. No.:</b> HY-N2450	Sunitinib (SU 11248)	<b>Cat. No</b> .: HY-10255A
Sulforaphene, isolated from radish seeds, exhibits an ED ₅₀ against velvetleaf seedlings approximately 2 x $10^{-4}$ M. Sulforaphene promotes cancer cells apoptosis and inhibits migration via inhibiting EGFR, p-ERK1/2, NF $\kappa$ B and other signals.	S ₂ C ₂ N N	Sunitinib (SU 11248) is a multi-targeted receptor tyrosine kinase inhibitor with $IC_{50}$ s of 80 nM and 2 nM for VEGFR2 and PDGFR $\beta$ , respectively.	Contraction of the second seco
Purity:99.26%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg		Purity:         98.96%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 100 mg, 200 mg, 500 mg	
Sunitinih Malate		Sunitinih-d10	
(SU 11248 Malate)	Cat. No.: HY-10255	(SU 11248-d10)	Cat. No.: HY-10255AS
Sunitinib Malate (SU 11248 Malate) is a multi-targeted receptor tyrosine kinase inhibitor with $IC_{50}$ S of 80 nM and 2 nM for VEGFR2 and PDGFR $\beta$ , respectively.	Contraction of the second s	Sunitinib D10 (SU 11248 D10) is a deuterium labeled Sunitinib. Sunitinib is a multi-targeted receptor tyrosine kinase inhibitor with $IC_{so}$ s of 80 nM and 2 nM for VEGFR2 and PDGFR $\beta$ , respectively.	
Purity:         99.47%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 100 mg, 200 mg, 500 mg		Purity:99.89%Clinical Data:No Development ReportedSize:1 mg, 5 mg	21
Sunitinib-d4	<b>Cat. No.:</b> HY-10255AS1	Supinoxin (RX-5902)	Cat. No.: HY-123611
Sunitinib-d4 (SU 11248-d4) is the deuterium labeled Sunitinib. Sunitinib (SU 11248) is a multi-targeted receptor tyrosine kinase inhibitor with $IC_{50}$ of 80 nM and 2 nM for VEGFR2 and PDGFR $\beta$ , respectively.		Supinoxin (RX-5902) is an orally active inhibitor of <b>phosphorylated-p68 RNA helicase (P-p68)</b> and a potent first-in-class <b>anti-cancer agent</b> . Supinoxin interacts with Y593 phosphorylated-p68 and attenuates the nuclear shuttling of β-catenin.	PCT NTA PT NTA PT NTA
Purity:>98%Clinical Data:Size:2.5 mg, 1 mg, 25 mg		Purity:99.90%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg	0.,

Suramin		Suramin sodium salt	
	Cat. No.: HY-B0879	(Suramin hexasodium salt)	Cat. No.: HY-B0879A
$\label{eq:suramin} \begin{array}{llllllllllllllllllllllllllllllllllll$	West West Strate		
SW106065		SY-5609	
	Cat. No.: HY-124778	(CDK7-IN-3)	Cat. No.: HY-138293
SW106065 is an <b>apoptosis</b> inducer in malignant peripheral nerve sheath tumors (MPNST). SW106065 inhibits ATP consumption of sMPNST and other models of MPNST with an $EC_{so}$ of 1 µM. SW106065 can be used for MPNST research.Purity:99.78% Clinical Data:Clinical Data:No Development Reported Size:10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10		$\begin{array}{llllllllllllllllllllllllllllllllllll$	HZ H F F F F F F F F F F F F F F F F F F
C71 D1 41		T 1101 toculate	
52L F1-41	Cat. No.: HY-100237	(TAI-95 tosylate)	Cat. No.: HY-120356A
SZL P1-41 is a specific <b>Skp2</b> inhibitor, binds to the F-box domain of Skp2 to prevent Skp1 association and Skp2 SCF complex formation. SZL P1-41, like Skp2 deficiency, augments p27-mediated apoptosis/senescence, while it impairs Akt-driven glycolysis. Anti-tumor activities. <b>Purity:</b> ≥98.0%		T-1101 tosylate (TAI-95 tosylate) is a Hec1/Nek2 (Highly expressed in cancer 1 / NIMA-related kinase 2) inhibitor with antitumor activity. T-1101 tosylate is inactive toward normal cells, kinases and hERG. Purity: 99.61%	04990 the
Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	00 mg	Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
	<u> </u>		
T-2 Toxin		T0901317	
(T-2 Mycotoxin)	Cat. No.: HY-N6792		Cat. No.: HY-10626
T-2 Toxin (T-2 Mycotoxin) is a toxic trichothecene mycotoxin produced by various Fusarium species in feedstuffs and cereal grains, $LD_{50}$ values of T-2 Toxin in mice and rats are 5.2 and 1.5 mg/kg BWa,respectively .Purity: $\geq 99.0\%$ Clinical Data:No Development Reported Size:Size:5 mg, 10 mg, 50 mg, 100 mg	Llot tot of of our	$\label{eq:constraint} \begin{array}{ll} T0901317 \mbox{ is an orally active and highly selective} \\ LXR agonist with an EC_{so} of 20 nM for LXR\alpha. \\ T0901317 \mbox{ activates FXR with an EC_{so} of 5 $\mu$M}. \\ T0901317 \mbox{ is ROR} $\alpha$ and ROR$ dual inverse agonist with $K$ values of 132 nM and 51 nM, respectively. \\ \hline Purity: $9.91\% \\ \hline Clinical Data: No Development Reported \\ \hline Size: $10 mM $\times 1 mL$, 10 mg, 50 mg, 100 mg \\ \hline \end{array}$	F F F F F F F F F F F F F F F F F F F
Terrelandida A		To continue line	
i accaionolide A	<b>Cat. No.:</b> HY-N2416	(N-acetyldinaline; CI-994; Goe-5549)	<b>Cat. No.:</b> HY-50934
Taccalonolide A is a microtubule stabilizer, which is a steroid isolated from Tacca chantrieri, with cytotoxic and antimalarial activities. Taccalonolide A causes $G_2$ -M accumulation, Bcl-2 phosphorylation and initiation of apoptosis.		Tacedinaline (N-acetyldinaline) is an inhibitor of the histone deacetylase (HDAC) with IC _{so} s of 0.9, 0.9, 1.2 $\mu$ M for recombinant HDAC 1, 2 and 3 respectively.	N NH2
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg	o	Purity:         99.55%           Clinical Data:         Phase 3           Size:         10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg	g, 500 mg

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Tadalafil		TAI-1	
(IC-351)	Cat. No.: HY-90009A		Cat. No.: HY-B0790
Tadalafil (IC-351) is a PDE5 inhibitor with an $\mathrm{IC}_{\mathrm{s0}}$ value of 1.8 nM.	H N O	TAI-1, an orally active anticancer agent, is a highly potent first-in-class $Hec1$ inhibitor, with a $GI_{s0}$ of 13.48 nM in K562 cells.	~a.d.s.ho
Purity:99.86%Clinical Data:LaunchedSize:10 mM × 1 mL, 50 mg, 100 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
TAK-243 (MLN7243)	<b>Cat. No.:</b> HY-100487	Takinib (EDHS-206)	<b>Cat. No.</b> : HY-103490
TAK-243 (MLN7243) is a first-in-class, selective ubiquitin activating enzyme, <b>UAE (UBA1)</b> inhibitor ( $IC_{s0}$ =1 nM), which blocks ubiquitin conjugation, disrupting monoubiquitin signaling as well as global protein ubiquitination.		Takinib (EDHS-206) is an orally active and selective <b>TAK1</b> inhibitor ( $IC_{s0}$ =9.5 nM), more than 1.5 log more potent than the second and third ranked targets, IRAK4 (120 nM) and IRAK1 (390 nM), respectively.	
Purity:         98.38%           Clinical Data:         Phase 1           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	○ N 6 - 5 - 55-331	Purity:99.15%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
l alampanel (GYKI-53773; LY-300164)	<b>Cat. No.:</b> HY-15079	laltobulin (HTI-286; SPA-110)	<b>Cat. No.:</b> HY-15584
Talampanel (LY300164) is an orally and selective $\alpha$ -amino-3-hydroxy-5-methyl-4-isoxazolepropionate (AMPA) receptor antagonis with anti-seizure activity. Talampanel (IVAX) has neuroprotective effects in rodent stroke models.	LN HTO	Taltobulin (HTI-286), a synthetic analogue of the tripeptide hemiasterlin, is a potent <b>antimicrotubule</b> agent that circumvents P-glycoprotein-mediated resistance in vitro and in vivo.	MH NH
Purity:         98.02%           Clinical Data:         Phase 2           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	NH ₂	Purity:99.90%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg	
Taltabulin bydrachlarida		Taltabulin trifluoroacatata	
(HTI-286 hydrochloride; SPA-110 hydrochloride)	Cat. No.: HY-15584B	(HTI-286 trifluoroacetate; SPA-110 trifluoroacetate)	Cat. No.: HY-15584A
Taltobulin hydrochloride (HTI-286 hydrochloride),         a synthetic analogue of the tripeptide         hemiasterlin, is a potent antimicrotubule agent         that circumvents P-glycoprotein-mediated         resistance in vitro and in vivo.         Purity:       98.34%         Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 5 mg, 10 mg, 50 mg	NH NH H-CI	Taltobulin trifluoroacetate (HTI-286         trifluoroacetate), a synthetic analogue of the         tripeptide hemiasterlin, is a potent         antimicrotubule agent that circumvents         P-glycoprotein-mediated resistance in vitro and in         vivo.         Purity:       99.96%         Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 5 mg, 10 mg	
Tamibarotene (Am 80)	<b>Cat. No.:</b> HY-14652	Tamoxifen (ICI 47699; (Z)-Tamoxifen; trans-Tamoxifen)	<b>Cat. No.:</b> HY-13757A
Tamibarotene is a <b>retinoic acid receptor</b> $\alpha/\beta$ (RAR $\alpha/\beta$ ) agonist, showing high selectivity over RAR $\gamma$ .		Tamoxifen (ICI 47699) is an orally active, selective <b>estrogen receptor</b> modulator ( <b>SERM</b> ) which blocks estrogen action in breast cells and can activate estrogen activity in other cells, such as bone, liver, and uterine cells.	N-00
Purity:         99.94%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 10 mg, 50 mg		Purity:         99.92%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 500 mg, 1 g, 5 g	~



Targaprimir-96 TFA		TAS-117	
	Cat. No.: HY-135276A		Cat. No.: HY-19934
Targaprimir-96 TFA is a potent inhibitor ofmicroRNA-96 (miR-96) processing. Targaprimir-96TFA selectively modulates miR-96 production incancer cells and triggers apoptosis. Targaprimir-96TFA binds primary miR-96 (pri-miR-96) with lownanomolar affinity.Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	night night of	TAS-117 is a potent, selective, orally active allosteric Akt inhibitor (with IC ₅₉ s of 4.8, 1.6, and 44 nM for Akt1, 2, and 3, respectively).TAS-117 triggers anti-myeloma activities and enhances fatal endoplasmic reticulum (ER) stress induced by proteasome inhibition.Purity:>98%Clinical Data:No Development Reported Size:5 mg, 10 mg, 25 mg, 50 mg	
TAS-117 hydrochloride	<b>Cat. No.</b> : HY-19934A	TAS6417 (CLN-081)	Cat. No.: HY-112299
TAS-117 hydrochloride is a potent, selective, orally active allosteric <b>Akt</b> inhibitor (with <b>IC</b> _{sp5} of 4.8, 1.6, and 44 nM for Akt1, 2, and 3, respectively). <b>Purity:</b> 98.96%		TAS6417 (CLN-081) is a highly effective, orally active and pan-mutation-selective EGFR tyrosine kinase inhibitor with a unique scaffold fitting into the ATP-binding site of the EGFR hinge region, with IC ₅₀ values ranging from 1.1-8.0 nM.Purity:98.77%	
Clinical Data: Phase 2		Clinical Data: Phase 2	100 mg
Size: 5 mg, 10 mg, 25 mg		Size: 10 mivi × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	luu mg
Tasisulam (LY 573636)	<b>Cat. No.:</b> HY-14804	Tasisulam sodium (LY 573636 sodium)	<b>Cat. No.</b> : HY-14804A
Tasisulam is a anticancer agent and induces <b>apoptosis</b> via the intrinsic pathway, resulting in cytochrome c release and caspase-dependent cell death. Tasisulam inhibits mitotic progression and induces vascular normalization.	CI C	Tasisulam is a anticancer agent and induces <b>apoptosis</b> via the intrinsic pathway, resulting in cytochrome c release and caspase-dependent cell death. Tasisulam inhibits mitotic progression and induces vascular normalization.	CI C
Purity:99.80%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg, 25 mg, 50 mg		Purity:>98%Clinical Data:Phase 3Size:1 mg, 5 mg	
Taurachanadaawychalic acid		Taurashanadaawushalis asid sadium	
(12-Deoxycholyltaurine)	Cat No: HY-N2027	(12-Deoxycholyltaurine sodium)	Cat No: HY-N1429
Taurochenodeoxycholic acid (12-Deoxycholyltaurine) is one of the main bioactive substances of animals' bile acid. Taurochenodeoxycholic acid induces <b>apoptosis</b> and shows obvious anti-inflammatory and immune regulation properties.	ности на стан	Taurochenodeoxycholic acid (12-Deoxycholyltaurine) sodium is one of the main bioactive substances of animals' bile acid. Taurochenodeoxycholic acid sodium induces <b>apoptosis</b> and shows obvious anti-inflammatory and immune regulation properties.	HOL H CH
Purity: 99.80% Clinical Data: Launched		Purity: ≥98.0% Clinical Data: Launched	
Size: 10 mM × 1 mL, 25 mg, 50 mg, 100 mg		Size: 100 mg	
Taurochenodeoxycholic acid-d4 sodium		Taurochenodeoxycholic acid-d4-1 sodium	
(12-Deoxycholyltaurine-d4 sodium)	Cat. No.: HY-N2027S	(12-Deoxycholyltaurine-d4-1 sodium)	Cat. No.: HY-N1429S2
Taurochenodeoxycholic acid-d4 (12-Deoxycholyltaurine-d4) sodium is the deuterium labeled Taurochenodeoxycholic acid. Taurochenodeoxycholic acid (12-Deoxycholyltaurine) is one of the main bioactive substances of animals' bile acid.		Taurochenodeoxycholic acid-d4-1 (sodium) is the deuterium labeled Taurochenodeoxycholic acid. Taurochenodeoxycholic acid (12-Deoxycholyltaurine) sodium is one of the main bioactive substances of animals' bile acid.	
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg, 10 mg		Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	

Taurochenodeoxycholic acid-d5 sodium		Taurodeoxycholic acid sodium hydrate	
(12-Deoxycholyltaurine-d5 sodium)	Cat. No.: HY-N1429S1	(Sodium taurodeoxycholate monohydrate)	Cat. No.: HY-B1899A
Taurochenodeoxycholic acid-d5 (12-Deoxycholyltaurine-d5) sodium is the deuterium labeled Taurochenodeoxycholic acid sodium.		Taurodeoxycholic acid sodium hydrate (Sodium taurodeoxycholate monohydrate) prevents apoptosis by blocking a calcium-mediated apoptotic pathway as well as caspase-12 activation.	HOCH THE HO OH CHE THE HO
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:         ≥98.0%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 10 mg, 50 mg	
Taurolidine	Cat. No.: HY-W011522	Tauroursodeoxycholate (Tauroursodeoxycholic acid; TUDCA; UR 906)	<b>Cat. No.:</b> HY-19696
Taurolidine is a broad-spectrum <b>antimicrobial</b> for the prevention of central venous catheter-related infections. Taurolidine has a direct and selective antineoplastic effect on brain tumor cells by the induction of <b>apoptosis</b> .		Tauroursodeoxycholate (Tauroursodeoxycholic acid) is an endoplasmic reticulum (ER) stress inhibitor. Tauroursodeoxycholate significantly reduces expression of apoptosis molecules, such as <b>caspase-3</b> and <b>caspase-12</b> . Tauroursodeoxycholate also inhibits <b>ERK</b> .	HOL CH CH CH
Purity:         ≥95.0%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 10 mg, 50 mg, 100 mg		Purity:     ≥ 98.0%       Clinical Data:     No Development Reported       Size:     10 mM × 1 mL, 50 mg	
Tauroursodeoxycholate dihydrate (Tauroursodeo dihydrate; TUDCA dihydrate; UR 906 dihydrate)	xycholic acid Cat. No.: HY-19696B	Tauroursodeoxycholate sodium (Tauroursodeoxyc sodium; TUDCA sodium; UR 906 sodium)	holic acid Cat. No.: HY-19696A
Tauroursodeoxycholate (Tauroursodeoxycholic acid; TDUCA) dihydrate is an endoplasmic reticulum (ER) stress inhibitor. Tauroursodeoxycholate significantly reduces expression of apoptosis molecules, such as caspase-3 and caspase-12. Tauroursodeoxycholate also inhibits ERK.	HO (1 H HO HO (1 H HO (1 H HO HO (1 H HO HO (1 H HO HO (1 H HO (1 H HO HO (1	Tauroursodeoxycholate (Tauroursodeoxycholic acid; TUDCA) sodium is an endoplasmic reticulum (ER) stress inhibitor. Tauroursodeoxycholate significantly reduces expression of apoptosis molecules, such as <b>caspase-3</b> and <b>caspase-12</b> . Tauroursodeoxycholate also inhibits <b>ERK</b> .	
Purity:     ≥98.0%       Clinical Data:     No Development Reported       Size:     10 mM × 1 mL, 50 mg		Purity:98.63%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 100 mg, 500 mg	
Tauroursodeoxycholate-d4		Tauroursodeoxycholate-d4 sodium (Tauroursodeo	oxycholic acid-d4
(Tauroursodeoxycholic acid-d4; TUDCA-d4; UR 906-d4)	Cat. No.: HY-19696S1	sodium; TUDCA-d4 sodium; UR 906-d4 sodium)	Cat. No.: HY-19696AS
Tauroursodeoxycholate-d4 is deuterium labeled Tauroursodeoxycholate. Tauroursodeoxycholate (Tauroursodeoxycholic acid) is an endoplasmic reticulum (ER) stress inhibitor.	артан артан артан артан артан артан	Tauroursodeoxycholate-d4 (Tauroursodeoxycholic acid-d4) sodium is the deuterium labeled Tauroursodeoxycholate sodium. Tauroursodeoxycholate (Tauroursodeoxycholic acid; TUDCA) sodium is an endoplasmic reticulum (ER) stress inhibitor.	
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
Tauroursodeoxycholate-d4-1 (Tauroursodeoxycholic acid-d4-1; TUDCA-d4-1; UR 906-d	<b>4-1țat. No.:</b> HY-19696S2	Tauroursodeoxycholate-d5	<b>Cat. No.:</b> HY-19696S
Tauroursodeoxycholate-d4-1 is the deuterium labeled Tauroursodeoxycholate. Tauroursodeoxycholate (Tauroursodeoxycholic acid) is an endoplasmic reticulum (ER) stress inhibitor.		Tauroursodeoxycholate-d5 is the deuterium labeled Tauroursodeoxycholate. Tauroursodeoxycholate (Tauroursodeoxycholic acid) is an endoplasmic reticulum (ER) stress inhibitor.	$[0] \begin{bmatrix} 0 & 0 & 0 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \\ 0 & 0 &$
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg, 10 mg	



Temozolomide-d3	<b>Cat. No.</b> : HY-17364S	Temsirolimus (CCI-779)	<b>Cat. No.:</b> HY-50910
Temozolomide-d3 (NSC 362856-d3) is the deuterium labeled Temozolomide. Temozolomide (NSC 362856) is an oral active <b>DNA alkylating</b> agent that crosses the blood-brain barrier. Temozolomide is also a <b>proautophagic</b> and <b>proapoptotic</b> agent.		Temsirolimus is an inhibitor of <b>mTOR</b> with an $IC_{50}$ of 1.76 $\mu$ M. Temsirolimus activates <b>autophagy</b> and prevents deterioration of cardiac function in animal model.	J. J. J. Conce
Purity:>98%Clinical Data:Size:1 mg, 5 mg	02 1112	Purity:         99.56%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 100 mg	~~ ·
Temsirolimus-d3 (CCI-779-d3)	<b>Cat. No</b> .: HY-50910S	Temsirolimus-d3-1 (CCI-779-d3-1)	<b>Cat. No.</b> : HY-50910S2
Temsirolimus-d3 (CCI-779-d3) is the deuterium labeled Temsirolimus. Temsirolimus is an inhibitor of <b>mTOR</b> with an <b>IC</b> ₅₀ of 1.76 $\mu$ M. Temsirolimus activates <b>autophagy</b> and prevents deterioration of cardiac function in animal model.		Temsirolimus-d3-1 (CCI-779-d3-1) is the deuterium labeled Temsirolimus. Temsirolimus is an inhibitor of <b>mTOR</b> with an IC ₅₀ of 1.76 $\mu$ M. Temsirolimus activates <b>autophagy</b> and prevents deterioration of cardiac function in animal model.	
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg		Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	
l emsirolimus-d /		lertenadine	Cat No. LIV P1102
Temsirolimus-d7 (CCI-779-d7) is the deuterium labeled Temsirolimus. Temsirolimus is an inhibitor of mTOR with an $IC_{so}$ of 1.76 µM. Temsirolimus activates autophagy and prevents deterioration of cardiac function in animal model.Purity:>98%		Terfenadine ((±)-Terfenadine) is a potent         open-channel blocker of hERG with an IC ₅₀ of 204         nM. Terfenadine, an H1 histamine receptor         antagonist, acts as a potent apoptosis inducer in         melanoma cells through modulation of Ca ²⁺ homeostasis.         Purity:       99.93%	
Clinical Data:No Development ReportedSize:1 mg, 5 mg		Clinical Data:LaunchedSize:10 mM × 1 mL, 100 mg	
Terferedine d10		Terforenting all	
((+)-Terfenadine-d10: MDL-991-d10)	Cat No: HY-B119351	Terrenadine-d5	Cat No: HV-B11935
Terfenadine-d10 ((±)-Terfenadine-d10) is the deuterium labeled Terfenadine. Terfenadine (((±)-Terfenadine) is a potent open-channel blocker of hERG with an IC ₅₀ of 204 nM.		Terfenadine-d3 (( $\pm$ )-Terfenadine-d3) is the deuterium labeled Terfenadine. Terfenadine (( $\pm$ )-Terfenadine) is a potent open-channel blocker of <b>hERG</b> with an <b>IC</b> ₅₀ of 204 nM.	
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	1	Purity:>98%Clinical Data:No Development ReportedSize:2000 μg, 5 mg, 10 mg, 25 mg	7.02
Terrein	<b>Cat. No.</b> : HY-119808	Terrestrosin D	<b>Cat. No.:</b> HY-N5074
Terrein is a melanogenesis inhibitor. Terrein induces <b>apoptosis</b> in breast cancer cell lines . Terrein is an inhibitor of quorum sensing and c-di-GMP in Pseudomonas aeruginosa.	HQ OH	Terrestrosin D, a steroidal saponin from Tribulus terrestris L., induces cell cycle arrest and cancer cells apoptosis. Terrestrosin D has antiangiogenic activities.	ڰ ڰڹ ڰڹؿۊڮۊڮڎ
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg	- ~ `	Purity:98.83%Clinical Data:No Development ReportedSize:5 mg, 10 mg	

Tetrahydropalmatine		Tetrahydroxyquinone	DRO) Cat. No : HV P1106
Tetrahydropalmatine possesses analgesic effects.         Tetrahydropalmatine acts through inhibition of amygdaloid release of dopamine to inhibit an epileptic attack in rats.         Purity:       99.16%         Clinical Data:       Launched         Size:       10 mM × 1 mL, 10 mg, 50 mg		Tetrahydroxyquinone         (Tetrahydroxy-1,4-benzoquinone), a primitive         anticataract agent, is a redox active         benzoquinone. Tetrahydroxyquinone can take part in         a redox cycle with semiquinone radicals, leading         to the formation of reactive oxygen species (ROS).         Purity:       ≥95.0%         Clinical Data:       No Development Reported         Size:       10 mM × 1 mL 100 mg	
Tetrahydroxyquinone monohydrate	Cat. No.: HY-B1106A	Tetramethylcurcumin (FLLL31)	Cat. No.: HY-N2521
Tetrahydroxyquinone monohydrate (Tetrahydroxy-1,4-benzoquinone monohydrate), a primitive anticataract agent, is a redox active benzoquinone.         Purity:       ≥97.0%		Tetramethylcurcumin (FLLL31), derived from curcumin, specifically suppresses the phosphorylation of <b>STAT3</b> by binding selectively to Janus kinase 2 and the STAT3 Src homology-2 domain. Tetramethylcurcumin exhibits anti-inflammatory and anti-cancer effects. <b>Purity:</b> 99.91%	- type
Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 100 mg		Clinical Data:       No Development Reported         Size:       5 mg, 10 mg	
Tezacitabine	<b>Cat. No.:</b> HY-106014	TG101209	<b>Cat. No.:</b> HY-10410
Tezacitabine is a cytostatic and cytotoxic antimetabolite and a <b>nucleoside analogue</b> . Tezacitabine irreversibly inhibits the <b>ribonucleotide reductase</b> and interferes with <b>DNA</b> <b>replication</b> and <b>repair</b> . Tezacitabine effectively induces cells <b>apoptotic</b> .		TG101209 is a selective JAK2 inhibitor with $IC_{so}$ of 6 nM, less potent to Flt3 and RET with $IC_{so}$ of 25 nM and 17 nM, appr 30-fold selective for JAK2 than JAK3, and sensitive to JAK2V617F and MPLW515L/K mutations.	10 stroft
Purity:         99.32%           Clinical Data:         Phase 2           Size:         5 mg, 10 mg, 25 mg, 50 mg		Purity:         99.72%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
TH1834	<b>Cat No</b> : HY-123604	TH1834 dihydrochloride	Cat. No : HY-123604A
TH1834 is a specific <b>Tip60 (KAT5) histone</b> acetyltransferase (HAT) inhibitor. TH1834 induces apoptosis and increases DNA damage in breast cancer. TH1834 does not affect the activity of related histone acetyltransferase MOF. Anticancer activity. <b>Purity:</b> 98.86% Clinical Data: No Development Reported	aon othe	TH1834 dihydrochloride is a specific <b>Tip60 (KAT5)</b> histone acetyltransferase inhibitor. TH1834 dihydrochloride induces apoptosis and increases DNA damage in breast cancer. TH1834 dihydrochloride does not affect the activity of related histone acetyltransferase MOF. Anticancer activity. Purity: 99.68% Clinical Data: No Development Reported	and the state
Size:         5 mg, 10 mg, 50 mg, 100 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	
Thailanstatin D	<b>Cat. No.:</b> HY-139104	Thalidomide D4	Cat. No.: HY-14658S
Thailanstatin D, an analogue of Thailanstatin A, is able to inhibit <b>AR-V7 gene splicing</b> by interfering the interaction between U2AF65 and SAP155 and preventing them from binding to polypyrimidine tract located between the branch point and the 3' splice site.	South Contraction	Thalidomide D4 is a deuterium labeled Thalidomide. Thalidomide inhibits <b>cereblon (CRBN)</b> , a part of the <b>cullin-4 E3 ubiquitin ligase</b> complex CUL4-RBX1-DDB1, with a $K_d$ of ~250 nM, and has immunomodulatory, anti-inflammatory and anti-angiogenic cancer properties.	
Purity:     >98%       Clinical Data:     No Development Reported       Size:     5 mg, 10 mg, 25 mg		Purity:     98.03%       Clinical Data:     No Development Reported       Size:     5 mg, 10 mg	

Thapsigargin	<b>Cat. No.:</b> HY-13433	Thiamine hydrochloride (Thiamine chloride hydroch Vitamin B1 hydrochloride)	loride; Cat. No.: HY-N0680
Thapsigargin, an <b>endoplasmic reticulum (ER)</b> <b>stress</b> inducer, is an inhibitor of microsomal <b>Ca²⁺-ATPase</b> . Thapsigargin efficiently inhibits coronavirus (HCoV-229E, MERS-CoV, SARS-CoV-2) replication in different cell types.		Thiamine hydrochloride (Thiamine chloride hydrochloride) is an essential micronutrient needed as a cofactor for many central metabolic enzymes.	
Purity:99.95%Clinical Data:No Development ReportedSize:1 mg, 5 mg, 10 mg, 25 mg, 50 mg		Purity:         99.99%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 100 mg, 1 g	0.148.5
Thiamine monochloride-C13 hydrochloride	<b>Cat. No.:</b> HY-N0680S	Thiamine-13C3 hydrochloride (Thiamine chloride-1 hydrochloride; Vitamin B1-13C3 hydrochloride)	L <b>3C3</b> Cat. No.: HY-N0680S3
Thiamine monochloride-C13 hydrochloride is the deuterium labeled Thiamine hydrochloride. Thiamine hydrochloride (Thiamine chloride hydrochloride) is an essential micronutrient needed as a cofactor for many central metabolic enzymes.	N	Thiamine-13C3 (Thiamine chloride-13C3) hydrochloride is the 13C-labeled Thiamine (hydrochloride). Thiamine hydrochloride (Thiamine chloride hydrochloride) is an essential micronutrient needed as a cofactor for many central metabolic enzymes.	СГ ¹³ СН ₃ N N NH ₂ S OH HCI
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg		Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	
Thiamine-d3 hydrochloride (Thiamine chloride-d3 hydrochloride; Vitamin B1-d3 hydrochloride)	<b>Cat. No.:</b> HY-N0680S1	Thiamine-d4 hydrochloride (Thiamine chloride-d4 hydrochloride; Vitamin B1-d4 hydrochloride)	<b>Cat. No.:</b> HY-N0680S2
Thiamine-d3 (Thiamine chloride-d3) hydrochloride is the deuterium labeled Thiamine hydrochloride. Thiamine hydrochloride (Thiamine chloride hydrochloride) is an essential micronutrient needed as a cofactor for many central metabolic enzymes. Purity: >98%	D D D Cr Hci	Thiamine-d4 (hydrochloride) is deuterium labeled Thiamine (hydrochloride). Thiamine hydrochloride (Thiamine chloride hydrochloride) is an essential micronutrient needed as a cofactor for many central metabolic enzymes. Purity: >98%	
Clinical Data: No Development Reported Size: 1 mg, 5 mg		Clinical Data: No Development Reported Size: 1 mg, 5 mg	
Thienopyridone	<b>Cat. No.:</b> HY-128153	Thiocolchicine	<b>Cat. No.</b> : HY-116852
Thienopyridone is a potent and selectivephosphatase of regenerating liver (PRL)phosphatase inhibitor with IC ₅₀ of 173 nM, 277nM and 128 nM for PRL-1, PRL-2, and PRL-3,respectively. Thienopyridone shows minimal effectson other phosphatases.Purity:98.04%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 50 mg, 100 mg	HN S S	Thiocolchicine, a derivative modified in the C Ring of Colchicine (HY-16569) with enhanced biological properties. Thiocolchicine is a potent inhibitor of <b>tubulin polymerization</b> ( $IC_{so}$ =2.5 $\mu$ M) and competitively binds to tubulin with a K _i of 0.7 $\mu$ M. Purity: >98% Clinical Data: No Development Reported Size: 1 mg, 5 mg	S S S S S S S S S S S S S S S S S S S
Thiocolchicine-d3	<b>Cat. No.:</b> HY-116852S	Thioridazine	<b>Cat. No.:</b> HY-B0965A
Thiocolchicine-d3 is deuterium labeled Thiocolchicine. Thiocolchicine, a derivative modified in the C Ring of Colchicine (HY-16569) with enhanced biological properties.	off off off NH	Thioridazine, an antagonist of the <b>dopamine</b> receptor D2 family proteins, exhibits potent anti-psychotic and anti-anxiety activities. Thioridazine is also a potent inhibitor of PI3K-Akt-mTOR signaling pathways with anti-angiogenic effect.	
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	u d d	Purity:>98%Clinical Data:LaunchedSize:1 mg, 5 mg	`\~_s^_"

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TM5441	<b>Cat. No.:</b> HY-101761	TMI-1 (WAY-171318)	<b>Cat. No.</b> : HY-101448
TM5441 is an orally bioavailable inhibitor of plasminogen activator inhibitor-1 (PAI-1), has IC ₅₀ values between 13.9 and 51.1 μM and induces intrinsic apoptosis in several human cancer cell lines.         Purity:       98.18%         Clinical Data:       No Development Reported         Size:       5 mg, 10 mg, 25 mg, 50 mg, 100 mg		TMI-1 is a potent inhibitor of disintegrin         metalloenzyme 17 (ADAM17) and other MMPs. TMI-1         inhibits LPS-induced TNF-α secretion in human         primary monocytes, and human whole blood.         Purity:       >98%         Clinical Data:       No Development Reported         Size:       1 mg, 5 mg	HO. HO. N.
Toddaculin	<b>Cat. No.:</b> HY-N9359	Tofacitinib (Tasocitinib; CP-690550)	<b>Cat. No.:</b> HY-40354
Toddaculin is a natural coumarin that can induce differentiation and apoptosis in leukemic cells.Toddaculin suppresses excess osteoclast activity and enhances osteoblast differentiation and mineralization. Toddaculin also exhibits anti-inflammatory activity.Purity:>98%Clinical Data:No Development ReportedSize:5 mg	of of of	Tofacitinib is an orally available JAK3/2/1 inhibitor with IC ₅₀ s of 1, 20, and 112 nM, respectively.         Purity:       99.99%         Clinical Data:       Launched         Size:       10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg	$ \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \begin{array}{c} \end{array}\\ \end{array}\\ \end{array}\\ \end{array}\\ \begin{array}{c} \end{array}\\ \end{array}\\ \end{array}\\ \begin{array}{c} \end{array}\\ \end{array}\\ \end{array}\\ \begin{array}{c} \end{array}\\ \end{array}\\ \end{array} $
Tofacitinih citrata		Tofacitinih Prodrug_1	
(Tasocitinib citrate; CP-690550 citrate)	Cat. No.: HY-40354A		Cat. No.: HY-145829
Tofacitinib citrate is an orally available         JAK1/2/3 inhibitor with IC ₅₀ s of 1, 20, and 112         nM, respectively. Tofacitinib citrate has         antibacterial, antifungal and antiviral         activities.         Purity:       99.98%         Clinical Data:       Launched		Tofacitinib Prodrug-1 is an effective and oral active prodrug to mitigate the systemic adverse effects of Tofacitinib. Tofacitinib Prodrug-1 can effectively attenuate the oxazolone-induced colitis in mice model with low toxicity. Purity: >98% Clinical Data: No Development Reported	J.C. Blendya.
Size: 10 mixi × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg	, 500 mg	Size: 1 mg, 5 mg	
Tofacitinib-13C3 (Tasocitinib-13C3; CP-690550-13C3)	<b>Cat. No.:</b> HY-40354S	Tofacitinib-d3 citrate (Tasocitinib-d3 citrate; CP-690550-d3 citrate)	<b>Cat. No.:</b> HY-40354AS
Tofacitinib-13C3 (Tasocitinib-13C3) is the 13C-labeled Tofacitinib. Tofacitinib is an orally available <b>JAK3/2/1</b> inhibitor with <b>IC</b> _{so} s of 1, 20, and 112 nM, respectively.	$\overset{^{\prime}}{\underset{N}{}} {\underset{N}{}} }{\underset{N}{}} {\underset{N}{}} {\underset{N}{}} }{\underset{N}{}} }{\underset{N}{}} {\underset{N}{}} }{\underset{N}{}} }{}} }{\underset{N}{}} }{}} }{\underset{N}{}} }{}} }{}{}} }{}{}} }{}}{}{}} }{}{}}}{}{}{}} }{}{}{}{}}{}{}}{}{}}{}{}}{}{}{}}{}{}{}}{}{}}{}{}{}}{}{}}{}{}{}{}}{}{}{}}{}{}{}{}{}}{}{}{}}{}{}{}}{}{}{}}{}{}{}{}{}{}{}{}}{}{}{}}{}{}{}{}{}{}}{}{}{}}{}{}{}}{}{}{}}{}{}{}{}{}{}{}{}{}{}{}{}{}{}}$	Tofacitinib-d3 (citrate) is deuterium labeled Tofacitinib (citrate). Tofacitinib citrate is an orally available JAK1/2/3 inhibitor with IC50s of 1, 20, and 112 nM, respectively. Tofacitinib citrate has antibacterial, antifungal and antiviral activities.	
Clinical Data:     No Development Reported       Size:     1 mg, 5 mg		Clinical Data: No Development Reported Size: 1 mg, 5 mg	
Tolcapone (Ro 40-7592)	<b>Cat. No.:</b> HY-17406	<b>Tolcapone-d4</b> (Ro 40-7592-d4)	<b>Cat. No.:</b> HY-17406S1
Tolcapone (Ro 40-7592) is a selective, orally active and powerful mixed (peripheral and central) <b>COMT</b> inhibitor with an IC ₅₀ of 773nM in the liver. Tolcapone is also a potent inhibitor of $\alpha$ -syn and A $\beta$ 42 oligomerization and fibrillogenesis.	HO HO O ^{-N[*]O[*]}	Tolcapone-d4 (Ro 40-7592-d4) is the deuterium labeled Tolcapone. Tolcapone (Ro 40-7592) is a selective, orally active and powerful mixed (peripheral and central) <b>COMT</b> inhibitor with an <b>IC</b> ₅₀ of 773nM in the liver.	
Purity:         99.74%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 10 mg, 50 mg, 100 mg	Land An	Purity:       >98%         Clinical Data:       No Development Reported         Size:       1 mg, 10 mg	







The effective of 4		Taslass	
(CS-045-d4)	Cat. No : HY-509355	TROIDX	Cat No: HY-101445
Troglitazone-d4 is deuterium labeled Troglitazone. Troglitazone is a PPARγ agonist, with EC50s of 550 nM and 780 nM for human and murinePPARγ receptor, respectively.		Trolox is an analogue of vitamin E with a powerful antioxidant effect. Trolox is also a powerful inhibitor of membrane damage.	но + + + + + + + + + + + + + + + + + + +
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg		Purity:99.87%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 500 mg, 1 g, 5 g	~
TrxR inhibitor D9	<b>Cat. No.:</b> HY-136279	TTNPB (Ro 13-7410; Arotinoid acid; AGN191183)	<b>Cat. No.:</b> HY-15682
TrxR inhibitor D9 is a potent and selective inhibitor of <b>thioredoxin reductase (TrxR)</b> , with an $EC_{s0}$ of 2.8 nM. TrxR inhibitor D9 has the capability to inhibit tumor proliferation both in vitro and in vivo.		TTNPB is a highly potent RAR agonist. Competitive binding assays using human RARs yield IC ₅₀ s of $\alpha$ =5.1 nM, $\beta$ = 4.5 nM, and $\gamma$ =9.3 nM, respectively.	XUL IN
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:98.81%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg	
Tubastatin A	<b>Cat. No.:</b> HY-13271A	Tubastatin A Hydrochloride (Tubastatin A HCl; TSA HCl)	<b>Cat. No.</b> : HY-13271
Tubastatin A is a potent and selective HDAC6 inhibitor with an IC _{so} of 15 nM in a cell-free assay, and is selective (1000-fold more) against all other isozymes except HDAC8 (57-fold more).         Purity:       98.12%         Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg	C R R OH	Tubastatin A (Hydrochloride) is a potent and selective HDAC6 inhibitor with IC ₅₀ of 15 nM in a cell-free assay, and is selective (1000-fold more) against all other isozymes except HDAC8 (57-fold more).         Purity:       98.21%         Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 2	200 mg
		Tubulin inhibitor 1	
(Tubeimoside-1; Lobatoside-H)	Cat. No.: HY-N0890		Cat. No.: HY-112607
Tubeimoside I(Lobatoside-H) is an extract from         Chinese herbal medicine Bolbostemma paniculatum         (MAXIM.) FRANQUET (Cucurbitaceae) has been shown         as a potent anti-tumor agent for a variety of         human cancers.         Purity:       99.70%         Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 10 mg, 50 mg		Tubulin inhibitor 1 is a tubulin inhibitor,         inhibits tubulin polymerization. Tubulin inhibitor         1 shows potent anti-tumor activity, casues         cellular mitotic arrest in the G2/M phase, and         induces cellular apoptosis.         Purity:       99.67%         Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	of N-
Tubulin inhibitor 17	<b>Cat. No.</b> : HY-144748	Tubulin inhibitor 23	<b>Cat. No.:</b> HY-144818
Tubulin inhibitor 17 (Compound 3b) is a <b>tubulin</b> <b>polymerization</b> inhibitor with an $IC_{so}$ of 12.38 $\mu$ M. Tubulin inhibitor 17 has anticancer activities and induces cell <b>apoptosis</b> .	N N	Tubulin inhibitor 23 is a potent <b>Tubulin</b> inhibitor with an <b>IC</b> _{so} of 4.8 $\mu$ M. Tubulin inhibitor 23 induces cell <b>apoptosis</b> . Tubulin inhibitor 23 shows antiangiogenic activity in a dose-dependent manner. Tubulin inhibitor 23 has the potential for the research of leukaemia.	
Purity:> 98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	N	Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~

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Tubulin polymerization-IN-17		Tubulin polymerization-IN-3	
	Cat. No.: HY-146362		Cat. No.: HY-145868
Tubulin polymerization-IN-17 (compound 23g) is a potent inhibitor of <b>tubulin</b> polymerization. Tubulin polymerization-IN-17 exhibits tubulin depolymerization and induced cell <b>apoptosis</b> and inhibits migration.		Tubulin polymerization-IN-3 (compound 4c) is a potent <b>tubulin polymerization</b> inhibitor, with an $IC_{so}$ of 3.84 $\mu$ M. Tubulin polymerization-IN-3 can induce apoptosis in colon cancer cells.	
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	~	Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	a
Tubulin polymerization-IN-4	<b>Cat. No.:</b> HY-144786	Tubulin polymerization-IN-5	<b>Cat. No.</b> : HY-144299
Tubulin polymerization-IN-4 is a potent <b>tubulin polymerization</b> inhibitor with $IC_{50}$ value of 4.6 $\mu$ M.		Tubulin polymerization-IN-5 (compound 20q) is a potent <b>tubulin</b> inhibitor with potential anticancer activities. Tubulin polymerization-IN-5 can arrest ESCC cells at G2/M phase and cause cells <b>apoptosis</b> .	And a land a
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg		Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	20 - 20
Tubulin polymerization-IN-6	<b>Cat. No.:</b> HY-146505	Tubulin polymerization-IN-9	<b>Cat. No.</b> : HY-146718
Tubulin polymerization-IN-6 (compound 5f) is a potent <b>tubulin polymerization</b> inhibitor, with an $IC_{so}$ of 1.09 $\mu$ M. Tubulin polymerization-IN-6 inhibits cell migration and tube formation and contributes to the anti-angiogenesis. <b>Purity:</b> >98% <b>Clinical Data:</b> No Development Reported		Tubulin polymerization-IN-9 is a potent <b>tubulin</b> inhibitor with IC ₅₀ of 1.82 μM. Tubulin polymerization-IN-9 causes cell cycle arrest at G2/M phase, and induces cell <b>apoptosis</b> and depolarized mitochondria of K562 cells. <b>Purity:</b> >98% <b>Clinical Data:</b> No Development Reported	
Size: 1 mg, 5 mg		Size: 1 mg, 5 mg	
Tubuloside B	<b>Cat. No.:</b> HY-N7637	Tubulysin A (TubA)	<b>Cat. No.:</b> HY-15995
Tubuloside B, one of the phenylethanoids isolated from the stems of Cistanche salsa, inhibits TNFα-induced apoptosis. Tubuloside B possesses antioxidative effects.		Tubulysin A(TubA) is a myxobacterial product that can function as an antiangiogenic agent in many in vitro assays; anti-microtubule, anti-mitotic, an apoptosis inducer, anticancer, anti-angiogenic, and antiproliferative.	
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg	LJ _{OH}	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
Tubulysin C	<b>Cat. No.:</b> HY-N2347	Tubulysin D	<b>Cat. No.:</b> HY-N2348
Tubulysin C is a highly cytotoxic peptide isolated from the myxobacterial species Archangium geophyra and Angiococcus disciformis.		Tubulysin D is one of the most potent derivatives among the tubulysins isolated from the myxobacterial species Archangium geophyra and Angiococcus disciformis.	
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg		Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg	

l ubulysin E		Tubulysin F	Cat No : HV-N7049
Tubulysin E is a highly cytotoxic peptide isolated from the myxobacterial species Archangium geophyra and Angiococcus disciformis.		Tubulysin F is a highly cytotoxic peptide isolated from the myxobacterial species Archangium geophyra and Angiococcus disciformis.	
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg		Purity:>98%Clinical Data:Size:5 mg, 10 mg, 25 mg	
Tubulysin G	Cat No · HY-N7050	Tubulysin H	Cat No · HY-N7051
Tubulysin G is a highly cytotoxic peptide isolated from the myxobacterial species Archangium geophyra and Angiococcus disciformis.		Tubulysin H is a highly cytotoxic peptide isolated from the myxobacterial species Archangium geophyra and Angiococcus disciformis.	₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽
Purity:>98%Clinical Data:Size:5 mg, 10 mg, 25 mg		Purity:>98%Clinical Data:Size:5 mg, 10 mg, 25 mg	
Tubulysin I		Tubulysin M	
Tubulysin I is a highly cytotoxic peptide isolated from the myxobacterial species Archangium geophyra and Angiococcus disciformis.		Tubulysin M is a highly cytotoxic peptide isolated from the myxobacterial species Archangium geophyra and Angiococcus disciformis.	
Purity:>98%Clinical Data:Size:5 mg, 10 mg, 25 mg		Purity:>98%Clinical Data:Size:25 mg, 50 mg, 100 mg	
TVB-3166	<b>Cat. No</b> .: HY-120394	Tyrphostin AG 879 (AG 879)	<b>Cat. No.:</b> HY-20878
TVB-3166 is an orally-available, reversible, and selective <b>fatty acid synthase</b> (FASN) inhibitor with $IC_{so}^{s}$ of 42 nM and 81 nM for biochemical FASN and cellular palmitate synthesis, respectively. TVB-3166 induces <b>apoptosis</b> , and inhibits in-vivo xenograft tumor growth.	HN-N HH-I TVB-3105	Tyrphostin AG 879 (AG 879) is a tyrosine kinase inhibitor that inhibits $TrKA$ phosphorylation (IC $_{\rm 50}$ of 10 $\mu$ M), but not TrKB and TrKC.	HO NH2
Purity:         99.76%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity:       99.54%         Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 3	100 mg
Tyrphostin AG1296 (AG1296)	<b>Cat. No.</b> : HY-13894	Ubiquitin Isopeptidase Inhibitor I, G5 (NSC144303)	<b>Cat. No.</b> : HY-100738
Tyrphostin AG1296 is a potent and selective inhibitor of <b>platelet-derived growth factor</b> receptor (PDGFR), with an IC ₅₀ of 0.8 $\mu$ M.		Ubiquitin Isopeptidase Inhibitor I, G5 (NSC 144303) is an apoptosome-independent <b>caspase</b> and <b>apoptosis</b> activator with $IC_{s0}$ values of 1.76 and 1.6 $\mu$ M in E1A and E1A/C9DN cells, respectively.	°°°°°°°°°°°°°°°°°°°°°°°°°°°°°°°°°°°°°°
Purity:         99.25%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	00 mg	Purity:≥90.0%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	



Urolithin C	<b>Cat. No.:</b> HY-135897	Ursonic acid (3-Ketoursolic acid)	<b>Cat. No.:</b> HY-N1486
Urolithin C, a gut-microbial metabolite of Ellagic acid, is a glucose-dependent activator of <b>insulin</b> <b>secretion</b> . Urolithin C is a <b>L-type Ca²⁺ channel</b> opener and enhances <b>Ca²⁺ influx</b> .	но он	Ursolic acid, a naturally occurring triterpenoid, induces the apoptosis of human cancer cells through multiple signaling pathways.	
Purity:99.66%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg, 50 mg, 100 mg	Ū	Purity:     ≥98.0%       Clinical Data:     No Development Reported       Size:     10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg	/ \H
USP7-IN-9	<b>Cat. No.:</b> HY-146887	Uvarigrin	<b>Cat. No.:</b> HY-N2632
USP7-IN-9 is a highly potent <b>ubiquitin-specific</b> <b>protease 7 (USP7)</b> inhibitor with an IC ₅₀ value of 40.8 nM. USP7-IN-9 can induce <b>apoptosis</b> and arrest cell progression at G0/G1 and S phases in RS4; 11 cells.		Uvarigrin, isolated from the roots of Uvaria calamistrata, induces tumor multidrug resistance cell apoptosis and triggers Caspase-9 activation.	
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg		Purity:>98%Clinical Data:No Development ReportedSize:500 μg, 1 mg, 2 mg	
UZH1	<b>Cat No</b> : HY-134673	UZH1a	<b>Cat No</b> : HY-134673A
UZH1 is a racemate of UZH1a and UZH1b. UZH1a is a potent and selective <b>METTL3</b> inhibitor, with an <b>IC</b> _{s0} of 280 nM. UZH1b (IC _{s0} =28 µM) is essentially inactive. UZH1 can be used for epitranscriptomic modulation of cellular processes. UZH1 has antitumor activity. <b>Purity:</b> 99.88% <b>Clinical Data:</b> No Development Reported <b>Size:</b> 5 mg, 10 mg, 25 mg, 50 mg, 100 mg		UZH1a is a potent and selective METTL3inhibitor, with an IC50 Of 280 nM. UZH1a can beused for epitranscriptomic modulation of cellularprocesses. UZH1a has antitumor activity. UZH1aalso can be used as a chemical probe for studyingMETTL3.Purity:98.86%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
Valepotriate		Valinomycin	
Valepotriate, isolated from Valeriana jatamansi Jones, has anti-epileptic and anti-cancer activities. Purity: 99.93% Clinical Data: No Development Reported Size: 5 mg 10 mg 20 mg		Valinomycin (NSC 122023), a cyclic depsipeptide antibiotic, act as a potassium selective ionophore. Valinomycin (NSC 122023) inhibits lymphocyte proliferation by its effects on the cell membrane, and induces apoptosis in CHO cells. Purity: 99.05% Clinical Data: No Development Reported Size: 10 mM × 1 mL 5 mg 10 mg	
Vandetanib (ZD6474)	<b>Cat. No.</b> : HY-10260	Vandetanib hydrochloride (ZD6474 hydrochloride)	<b>Cat. No.:</b> HY-10260B
Vandetanib (D6474) is a potent, orally active inhibitor of VEGFR2/KDR tyrosine kinase activity ( $IC_{50}$ =40 nM). Vandetanib also has activity versus the tyrosine kinase activity of VEGFR3/FLT4 ( $IC_{50}$ =110 nM) and EGFR/HER1 ( $IC_{50}$ =500 nM).	N O O N N O O O N N O N N N N N H	Vandetanib hydrochloride (D6474 hydrochloride) is a potent, orally active inhibitor of VEGFR2/KDR tyrosine kinase activity (IC ₅₀ =40 nM). Vandetanib hydrochloride also has activity versus the tyrosine kinase activity of VEGFR3/FLT4 (IC ₅₀ =110 nM) and EGFR/HER1 (IC ₅₀ =500 nM).	
Purity:         99.89%           Clinical Data:         Launched           Size:         10 mM × 1 mL, 10 mg, 25 mg, 50 mg, 100 mg	Br ~~ F	Purity:>98%Clinical Data:LaunchedSize:1 mg, 5 mg	H-CI

Vandetanib trifluoroacetate		Vandetanib-d4	Cat. No : HV-1026051
Vandetanib trifluoroacetate (D6474 trifluoroacetate) is a potent, orally active inhibitor of VEGFR2/KDR tyrosine kinase activity (IC ₅₀ =40 nM).		Vandetanib-d4 (ZD6474-d4) is the deuterium labeled Vandetanib. Vandetanib (ZD6474) is a potent, orally active inhibitor of <b>VEGFR2/KDR</b> tyrosine kinase activity ( $IC_{s0}$ =40 nM).	
Purity:>98%Clinical Data:LaunchedSize:1 mg, 5 mg	F Br F	Purity:         >98%           Clinical Data:	Br
Vandetanib-d6 (ZD6474-d6)	<b>Cat. No.</b> : HY-10260S	Vanillyl alcohol (p-(Hydroxymethyl)guaiacol)	<b>Cat. No.:</b> HY-N2067
Vandetanib-d6 (ZD6474-d6) is the deuterium labeled Vandetanib. Vandetanib (D6474) is a potent, orally active inhibitor of VEGFR2/KDR tyrosine kinase activity ( $IC_{50}$ =40 nM).		Vanillyl alcohol (p-(Hydroxymethyl)guaiacol), derived from vanillin, is a phenolic alcohol and is used as a flavoring agent in foods and beverages.	НО ОН
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	or s r	Purity:     99.68%       Clinical Data:     No Development Reported       Size:     10 mM × 1 mL, 100 mg	
VAS 3947	<b>Cat. No.</b> : HY-111447	Vatalanib dihydrochloride (PTK787 dihydrochloride dihydrochloride; ZK-222584 dihydrochloride)	; <b>CGP-797870</b> <b>Cat. No.:</b> HY-12018
VAS 3947, a specific <b>NADPH oxidase (NOX)</b> inhibitor, exerts a potent antiplatelet effect. VAS3947 induces <b>apoptosis</b> independently of anti-NOX activity, via UPR activation, mainly due to aggregation and misfolding of proteins.	N N N N N	Vatalanib dihydrochloride (PTK787 dihydrochloride) is an inhibitor of $\rm VEGFR2/KDR$ with $\rm IC_{50}$ of 37 nM.	
Purity:99.59%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg, 50 mg, 100 mg	D	Purity:         99.97%           Clinical Data:         Phase 3           Size:         10 mM × 1 mL, 10 mg, 50 mg, 100 mg	N
Vatalanib-d4 dihydrochloride	<b>Cat. No.:</b> HY-12018S	VDR agonist 1	<b>Cat. No.</b> : HY-114310
Vatalanib-d4 (PTK787-d4) dihydrochloride is the deuterium labeled Vatalanib dihydrochloride. Vatalanib (PTK787) dihydrochloride is an inhibitor of VEGFR2/KDR with IC _{s0} of 37 nM.		VDR agonist 1 (compound 28) is a nonsteroidal <b>Vitamin D receptor (VDR)</b> agonist, with an <b>IC</b> _{so} of 690 nM in MCF-7 cells.	F. H. K.
Purity:>98%Clinical Data:Size:1 mg, 10 mg	N	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	
VEGFR-2-IN-11	<b>Cat. No.:</b> HY-145856	VEGFR-2-IN-13	<b>Cat. No.</b> : HY-144754
VEGFR-2-IN-11 (Compound 8h) is a potent <b>VEGFR-2</b> inhibitor with an <b>IC</b> _{so} of 60.27 nM. VEGFR-2-IN-11 shows antitumor activity and induces cell <b>apoptosis</b> .		VEGFR-2-IN-13 (Compound 19a) is a potent VEGFR-2 inhibitor with an $IC_{so}$ of 3.4 nM. VEGFR-2-IN-13 disrupts the HepG2 cell cycle by arresting the G2/M phase and induces <b>apoptosis</b> .	
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	Br	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	



Verteporfin		Verubulin	
(CL 318952)	Cat. No.: HY-B0146	(MPC 6827)	Cat. No.: HY-14907
Verteporfin (CL 318952) is a photosensitizer for photodynamic therapy to eliminate the abnormal blood vessels in the eye associated with conditions such as age-related macular degeneration. Verteporfin is a YAP inhibitor which disrupts YAP-TEAD interactions. Purity: 99.58% Clinical Data: Launched		Verubulin (MPC-6827) is a microtubule-disrupting agent with potent and broad-spectrum in vitro and in vivo cytotoxic activities, and acts as a promising candidate for the treatment of multiple cancer types. Purity: 99.34% Clinical Data: Phase 2	
Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg	
Vesatolimod		VII-31	
(GS-9620)	Cat. No.: HY-15601		Cat. No.: HY-133558
Vesatolimod (GS-9620) is a potent, selective and orally active agonist of Toll-Like Receptor (TLR7) with an $EC_{50}$ of 291 nM.		VII-31 is a potent <b>NEDDylation</b> pathway activator to inhibit the tumor progression in vitro and in vivo. VII-31 induces <b>apoptosis</b> via intrinsic and extrinsic pathways.	
Purity: 99.90%	19112	Purity: 98.28%	
Clinical Data: Phase 2 Size: 10 mM × 1 mL, 5 ma, 10 ma, 25 ma, 50 ma, 10	0 ma	Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 ma, 10 ma, 25 ma, 50 ma, 1	.00 ma
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Vildagliptin		Vildagliptin dihydrate	
(LAF237; NVP-LAF 237)	Cat. No.: HY-14291	(LAF237 dihydrate; NVP-LAF 237 dihydrate)	Cat. No.: HY-14291A
Vildagliptin (LAF237) is a potent, stable, selective dipeptidyl peptidase IV ( <b>DPP-IV</b> ) inhibitor with an IC ₅₀ of 3.5 nM in human Caco-2 cells. Vildagliptin possesse excellent oral bioavailability and potent antihyperglycemic activity. <b>Purity:</b> 98.18%	HO NH O NN CN	Vildagliptin dihydrate (LAF237 dihydrate) is a potent, stable, selective dipeptidyl peptidase IV (DPP-IV) inhibitor with an IC ₅₀ of 3.5 nM in human Caco-2 cells. Vildagliptin dihydrate possesses excellent oral bioavailability and potent antihyperglycemic activity. Purity: >98%	$H_2O$ $H_2O$ $H_2O$
Clinical Data: Launched		Clinical Data: Launched	0.000
Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 500 mg		Size: 1 mg, 5 mg	
Vilde ellection d2			
viidagiiptin-d3 (LAF237-d3: NVP-LAF 237-d3)	Cat No: HV-142915	viidagiiptin-d7 (LAF237-d7: NVP-LAF 237-d7)	Cat No : HV-1429151
Vildagliptin-d3 (LAF237-d3) is the deuterium labeled Vildagliptin. Vildagliptin (LAF237) is a potent, stable, selective dipeptidyl peptidase IV (DPP-IV) inhibitor with an $IC_{50}$ of 3.5 nM in human Caco-2 cells.		Vildagliptin-d7 is deuterium labeled Vildagliptin. Vildagliptin (LAF237) is a potent, stable, selective dipeptidyl peptidase IV (DPP-IV) inhibitor with an IC50 of 3.5 nM in human Caco-2 cells.	
Purity: >98%	N	Purity: >98%	$\checkmark$
Clinical Data: No Development Reported		Clinical Data: No Development Reported	
<b>512e</b> . 500 μg, 5 mg		Size. I mg, 5 mg	
Vin-C01	<b>Cat. No.:</b> HY-147502	Vin-F03	<b>Cat. No.:</b> HY-147503
Vin-C01 is a potent <b>pancreatic </b> $\beta$ <b>-cells</b> protective agent with an <b>EC</b> _{s0} of 0.22 $\mu$ M. Vin-C01 effectively promotes $\beta$ -cell survival and protects $\beta$ -cells from STZ-induced <b>apoptosis</b> . Vin-C01 can be used for type 2 diabetes mellitus research.		Vin-F03 is a potent <b>pancreatic</b> $\beta$ -cells protective agent with an EC ₅₀ of 0.27 $\mu$ M. Vin-F03 effectively promotes $\beta$ -cell survival and protects $\beta$ -cells from STZ-induced <b>apoptosis</b> . Vin-F03 can be used for type 2 diabetes mellitus research.	N H
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	

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Vincristine sulfate (Leurocristine sulfate: NSC-67574		Vincristine-d3 sulfate (Leurocristine-d3 sulfate:	
sulfate; 22-Oxovincaleukoblastine sulfate)	Cat. No.: HY-N0488	NSC-67574-d3 sulfate;)	Cat. No.: HY-N0488S
Vincristine sulfate is an antitumor vinca alkaloid         which inhibits microtubule formation in mitotic         spindle, resulting in an arrest of dividing cells         at the metaphase stage. It binds to microtubule         with a K ₁ of 85 nM.         Purity:       99.81%         Clinical Data:       Launched         Size:       10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg		Vincristine-d3 (Leurocristine-d3) sulfate is the deuterium labeled Vincristine sulfate. Vincristine sulfate is an antitumor vinca alkaloid which inhibits <b>microtubule</b> formation in mitotic spindle, resulting in an arrest of dividing cells at the metaphase stage. <b>Purity:</b> >98% <b>Clinical Data:</b> No Development Reported <b>Size:</b> 1 mg, 10 mg	Strate with
Vincristine-d3-ester sulfate (Leurocristine-d3-ester s	ulfate;	Vincristine-d6 sulfate (Leurocristine-d6 sulfate;	
NSC-67574-d3-ester sulfate;)	Cat. No.: HY-N0488S1	NSC-67574-d6 sulfate;)	Cat. No.: HY-N0488S2
Vincristine-d3-ester (Leurocristine-d3-ester) sulfate is the deuterium labeled Vincristine sulfate. Vincristine sulfate is an antitumor vinca alkaloid which inhibits <b>microtubule</b> formation in mitotic spindle, resulting in an arrest of dividing cells at the metaphase stage. <b>Purity:</b> >98% <b>Clinical Data:</b> No Development Reported <b>Size:</b> 1 mg	арана	Vincristine-d6 (Leurocristine-d6) sulfate is the deuterium labeled Vincristine sulfate. Vincristine sulfate is an antitumor vinca alkaloid which inhibits <b>microtubule</b> formation in mitotic spindle, resulting in an arrest of dividing cells at the metaphase stage. <b>Purity:</b> >98% <b>Clinical Data:</b> No Development Reported <b>Size:</b> 1 mg, 5 mg	
Vietucortik		Vistucertik d2	
(AZD2014)	Cat. No.: HY-15247	(AZD2014-d3)	Cat. No.: HY-15247S
Vistusertib (AZD2014) is an ATP competitive <b>mTOR</b> inhibitor with an $IC_{so}$ of 2.81 nM. AZD2014 inhibits both <b>mTORC1</b> and <b>mTORC2</b> complexes.	C, H C L M M	Vistusertib-d3 (AZD2014-d3) is the deuterium labeled Vistusertib. Vistusertib (AZD2014) is an ATP competitive <b>mTOR</b> inhibitor with an IC ₅₀ of 2.81 nM. AZD2014 inhibits both <b>mTORC1</b> and <b>mTORC2</b> complexes.	Strong of
Purity:         98.21%           Clinical Data:         Phase 2           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 7	200 mg	Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg	
Vitamin K4		Volasertib	
(acetomenaphthone)	Cat. No.: HY-B1508	(BI 6727)	Cat. No.: HY-12137
Vitamin K4 is a chemically synthesized Vitamin K         which plays an important role in the normal blood         coagulation system.         Purity:       99.89%         Clinical Data:       Launched         Size:       10 mM × 1 mL, 200 mg, 1 g		Volasertib (BI 6727) is an orally active, highly potent and ATP-competitive Polo-like kinase 1 (PLK1) inhibitor with an IC ₅₀ of 0.87 nM. Volasertib inhibits PLK2 and PLK3 with IC ₅₀ s of 5 and 56 nM, respectively. Volasertib induces mitotic arrest and <b>apoptosis</b> .Purity:99.41% Clinical Data:Phase 3 Size:10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	La ofation
Volasertib trihydrochloride (BI 6727 trihydrochloride)	<b>Cat. No.:</b> HY-12137A	Voreloxin (SNS-595; Vosaroxin; AG 7352)	<b>Cat. No.:</b> HY-10534
Volasertib (BI 6727) trihydrochloride is an orally active, highly potent and ATP-competitive <b>Polo-like</b> <b>kinase 1 (PLK1)</b> inhibitor with an IC ₅₀ of 0.87 nM. Volasertib trihydrochloride inhibits PLK2 and PLK3 with IC ₅₀ s of 5 and 56 nM, respectively.	ано но но	Voreloxin (SNS-595; Vosaroxin; AG 7352) is a first-in-class <b>topoisomerase II</b> inhibitor that intercalates DNA and induces site-selective DNA DSB, G2 arrest, and apoptosis.	NH N N N N N N N N N N N N N N N N N N
Purity:     >98%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg		Purity:     >98%       Clinical Data:     Phase 3       Size:     1 mg, 5 mg	

Varelayin Hydrochlarida (SNS 505 Hydrochlarida)	locarovin	Vorinostat	
Hydrochloride; AG 7352 Hydrochloride)	Cat. No.: HY-16518	(SAHA; Suberoylanilide hydroxamic acid)	Cat. No.: HY-10221
Voreloxin Hydrochloride is a first-in-class topoisomerase II inhibitor that intercalates DNA and induces site-selective DNA DSB, G2 arrest, and apoptosis. Purity: 99.96% Clinical Data: Phase 3 Size: 5 mo 10 mo 50 mo		Vorinostat (SAHA) is a potent and orally active pan-inhibitor of HDAC1, HDAC2 and HDAC3 (Class I), HDAC6 and HDAC7 (Class II) and HDAC11 (Class IV), with ID ₅₀ values of 10 nM and 20 nM for HDAC1 and HDAC3, respectively. Vorinostat induces cell <b>apoptosis</b> . <b>Purity:</b> 99.90% Clinical Data: Launched Size: 10 mM x 1 mL 250 mg 500 mg 1 g 5 g	С 4, в он
Size. 5 mg, 10 mg, 50 mg		Size. 10 milli × 1 mil, 250 mg, 500 mg, 1 g, 5 g	
Vorinostat-d5 (SAHA-d5; Suberoylanilide hydroxamic acid-d5)	<b>Cat. No.:</b> HY-115412	VR23	<b>Cat. No.:</b> HY-18741
Vorinostat-d5 (SAHA-d5) is the deuterium labeled Vorinostat. Vorinostat (SAHA) is a potent and orally active pan-inhibitor of HDAC1, HDAC2 and HDAC3 (Class I), HDAC7 (Class II) and HDAC11 (Class IV), with ID ₅₀ values of 10 nM and 20 nM for HDAC1 and HDAC3, respectively.		VR23 is a small molecule that potently inhibits the activities of trypsin-like proteasomes ( $IC_{so}$ =1 nM), chymotrypsin-like proteasomes ( $IC_{so}$ =50-100 nM), and caspase-like proteasomes ( $IC_{so}$ =3 $\mu$ M).	
Purity:     ≥99.0%       Clinical Data:     No Development Reported       Size:     1 mg		Purity:       98.05%         Clinical Data:       No Development Reported         Size:       5 mg, 10 mg, 50 mg, 100 mg	°₃ _{N¢} o.
VS 8	Cat. No.: HY-143491	V1P50469	Cat. No.: HY-114162
VS 8 (Compound VS 8) is a potent, orally active VEGFR-2 inhibitor with significant <b>anti-angiogenic</b> effects. VS 8 induces cancer cell <b>apoptosis</b> and migration. VS 8 is active against CSCs (Cancer stem cells).		VTP50469 is a potent, highly selective and orally active <b>Menin-MLL interaction</b> inhibitor with a <b>K</b> , of 104 pM. VTP50469 has potently anti-leukemia activity.	
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	V 0	Purity:99.41%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	<u></u>
VTP50469 fumarate		W-7 hydrochloride	
	Cat. No.: HY-114162A		Cat. No.: HY-100912
VTP50469 fumarate is a potent, highly selective and orally active Menin-MLL interaction inhibitor with a K _i of 104 pM. VTP50469 fumarate has potently anti-leukemia activity.         Purity:       98.84%         Clinical Data:       No Development Reported	N N N N N N N N N N N N N N N N N N N	W-7 hydrochloride is a selective <b>calmodulin</b> antagonist. W-7 hydrochloride inhibits the <b>Ca²⁺-calmodulin</b> -dependent <b>phosphodiesterase</b> and <b>myosin light chain kinase</b> with <b>IC</b> ₅₀ values of 28 $\mu$ M and 51 $\mu$ M, respectively. W-7 hydrochloride induces <b>apoptosis</b> and has antitumor activity. <b>Purity:</b> 99.65% <b>Clinical Data:</b> No Development Reported	CI H-CI H-CI NH2
Size: 5 mg, 10 mg, 50 mg, 100 mg		Size: 10 mM × 1 mL, 25 mg, 50 mg	
W146	<b>Cat. No.</b> : HY-101395	W146 TFA	<b>Cat. No.:</b> HY-101395A
W146 is a selective antagonist of sphingosine-1-phosphate receptor 1 (S1PR1) with an EC ₅₀ value of 398 nM.	чето сорн	W146 TFA is a selective antagonist of sphingosine-1-phosphate receptor 1 (S1PR1) with an $EC_{50}$ value of 398 nM.	
Purity:     ≥99.0%       Clinical Data:     No Development Reported       Size:     500 μg		Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg, 10 mg	

Waltonitone		WDR5-IN-1	
	Cat. No.: HY-128366		Cat. No.: HY-133121
Waltonitone is a ursane-type pentacyclic triterpene isolated from Gentian waltonii Burkill. Waltonitone significantly inhibits hepatocellular carcinoma cells growth and induces <b>apoptosis</b> in vitro and in vivo.	И И И И И И И И И И И И И И И И И И И	WDR5-IN-1 is a potent and selective <b>WD repeat</b> <b>domain 5 (WDR5)</b> inhibitor, with a $K_d$ of <0.02 nM. WDR5-IN-1 inhibits MLL1 histone methyltransferase (HMT) activity with an IC ₅₀ of 2.2 nM.	,ápap
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 25 mg	° ∕\A	Purity:         98.71%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	.00 mg
Wedelolactone	Cat. No.: HY-N0551	WEHI-9625	Cat. No.: HY-128777
Wedelolactone, a natural product from Ecliptae herba, suppresses LPS-induced <b>caspase-11</b> expression by directly inhibiting the IKK Complex. Wedelolactone inhibits <b>5-lipoxygenase</b> ( <b>5-Lox</b> ) (IC ₅₀ ~2.5 µM) activity by an oxygen radical scavenging mechanism. <b>Purity:</b> 99.91%	HO J J J O HO J J J O HO O	WEHI-9625 is a tricyclic sulfone, first-in-class inhibitor of <b>apoptosis</b> with an EC ₅₀ of 69 nM. WEHI-9625 binds to VDAC2 and promotes its ability to inhibit <b>apoptosis</b> driven by mouse BAK. WEHI-9625 is completely inactive against both human BAK and the closely related apoptosis effector BAX. Purity: 99.05% Clinical Data: No Davelopment Paported	CC. CC
Size: 10 mM × 1 mL, 5 mg, 10 mg, 20 mg		Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	.00 mg
WHI-P154		WM-3835	
	Cat. No.: HY-13895		Cat. No.: HY-134901
WHI-P154 is a potent EGFR inhibitor, and also modestly blocks JAK3, with $IC_{so}s$ of 4 nM and 1.8 $\mu M,$ respectively.	HN N	WM-3835 is a potent and high-specific <b>HBO1</b> ( <b>KAT7</b> or <b>MYST2</b> ) inhibitor and binds directly to the acetyl-CoA binding site of HBO1 33. WM-3835 activates <b>apoptosis</b> while inhibits osteosarcoma (OS) cell proliferation, migration and invasion.	
Purity:98.92%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 10 mg, 50 mg	o V N	Purity:98.10%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	.00 mg
		WP1000	
wogonin	Cat. No. LIV. NO.400	WP1066	Cot No. UV 15212
Wogonin is a naturally occurring mono-flavonoid,	Cat. No.: HY-N0400	WP1066 is an inhibitor of JAK2 and STAT3, and	Cat. No.: HY-15312
exhibits anti-inflammatory and anti-tumor effects.	HOLO	affecting JAK1 and JAK3.	
Purity: 99.98%	OH U	Purity: 99.90%	
Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg		Clinical Data:Phase 1Size:10 mM × 1 mL, 10 mg, 50 mg	
WT-161	Cat. No.: HY-100871	WYC-209	<b>Cat. No.:</b> HY-124136
WT-161 is a potent and selective HDAC6 inhibitor with an $IC_{_{50}}$ of 0.40 nM.	^{مو} لاً میں اور می	WYC-209, a synthetic retinoid, is a <b>retinoic acid</b> <b>receptor (RAR)</b> agonist. WYC-209 induces apoptosis primarily via the <b>caspase 3</b> pathway ( $(C_{so}=0.19\mu M$ for inmalignant murine melanoma TRCs), and has long-term effects with little toxicity.	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
Purity:         98.52%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	00 mg	Purity:99.64%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 50 mg, 100 mg	24

WYE-132	Cat. No : HV-10044	WYE-354	<b>Cat No</b> : HV-12034
WYE-123102)         WYE-132 (WYE-125132) is a highly potent,         ATP-competitive, and specific mTOR kinase         inhibitor (IC ₅₀ : 0.19±0.07 nM; >5,000-fold         selective versus PI3Ks). WYE-132 (WYE-125132)         inhibits mTORC1 and mTORC2.         Purity:       99.40%         Clinical Data:       No Development Reported         Size:       10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 50 mg, 100		$\label{eq:WYE-354} \begin{array}{ll} \text{WYE-354 is an ATP-competitive mTOR inhibitor} \\ \text{with an IC}_{s_0} \mbox{ of 5 nM. WYE-354 also inhibits PI3K} \\ \text{and PI3K} \mbox{ with IC}_{s_0} \mbox{ of 1.89 } \mu \mbox{M and 7.37 } \mu \mbox{M}, \\ \text{respectively. WYE-354 inhibits both mTORC1 and} \\ \text{mTORC2. WYE-354 induces autophagy activation in} \\ \text{vitro.} \\ \begin{array}{ll} \text{Purity:} & 98.0\% \\ \mbox{Clinical Data:} & \text{No Development Reported} \\ \mbox{Size:} & 10 \ \mbox{mM} \times 1 \ \mbox{mL}, \ \mbox{5 mg}, \ \mbox{20 mg}, \ \mbox{50 mg}, \ \mbo$	
Xanthatin	<b>Cat. No.:</b> HY-N3032	Xanthoangelol	<b>Cat. No.</b> : HY-111588
Xanthatin is isolated from Xanthium strumarium leaves. Purity: 99.79%	in the second	Xanthoangelol, extracted from Angelica keiskei, suppresses obesity-induced inflammatory responses. Xanthoangelol possesses antibacterial activity. Xanthoangelol inhibits monoamine oxidases. Xanthoangelol induces apoptosis in neuroblastoma and leukemia cells. Purity: 98.36%	HO,
Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Clinical Data: No Development Reported Size: 1 mg	
Xanthohumol	<b>Cat. No.:</b> HY-N1067	Xanthone	<b>Cat. No.:</b> HY-N0126
Xanthohumol is one of the principal flavonoids isolated from hops, the inhibitor of diacylglycerol acetyltransferase (DGAT), COX-1 and COX-2, and shows anti-cancer and anti-angiogenic activities.	HOTO OH	Xanthone is isolated from Mangosteen and is known to control cell division and growth, apoptosis, inflammation, and metastasis in different stages of carcinogenesis.	
Purity:         99.84%           Clinical Data:         Phase 1           Size:         10 mM × 1 mL, 5 mg, 10 mg, 25 mg		Purity:99.66%Clinical Data:No Development ReportedSize:100 mg	∞ .0. ∞
Xanthurenic acid	Cat. No.: HY-W014666	Xestospongin C ((-)-Xestospongin C)	<b>Cat. No.:</b> HY-103312
Xanthurenic acid is a putative endogenous <b>Group</b> II metabotropic glutamate receptor agonist, on sensory transmission in the thalamus.	ОН ОН	Xestospongin C ((-)-Xestospongin C) is a selective, reversible <b>inositol 1,4,5-trisphosphate</b> <b>receptor (IP3R)</b> inhibitor. Xestospongin C acts as an inhibitor of the sarcoplasmic/endoplasmic reticulum Ca ²⁺ ATPase (SERCA) pump of internal	
Purity:99.87%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 100 mg	о́н	Purity:       >98%         Clinical Data:       No Development Reported         Size:       10 µg, 25 µg	↓ H ⊂
XL019	<b>Cat. No.</b> : HY-13775	XMU-MP-3	<b>Cat. No.</b> : HY-136531
XL019 is a potent, orally active, and selective <b>JAK2</b> inhibitor, with <b>IC</b> ₅₀ s of 2.2, 134.3, and 214.2 nM for JAK2, JAK1 and JAK3, respectively.	0042046	XMU-MP-3 is a potent non-covalent <b>BTK</b> inhibitor with IC ₅₀ s of 10.7 nM and 17.0 nM for BTK WT and BTK C481S mutation in the presence of 10 $\mu$ M ATP, respectively. XMU-MP-3 also induces <b>apoptosis</b> .	-stopptor
Purity:         ≥98.0%           Clinical Data:         Phase 1           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity:98.27%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 50 mg, 100 mg	

XPO1-IN-1	<b>Cat. No.:</b> HY-144763	Xylopine	<b>Cat. No.:</b> HY-N9534
XPO1-IN-1 (compound D4) is an orally active and potent XPO1 inhibitor, with an IC ₅₀ of 24 nM in MM.1S cell. XPO1-IN-1 can efficiently induce cell apoptosis and cell cycle arrest. XPO1-IN-1 displays favorable metabolic stability and pharmacokinetic properties.Purity:>98%Clinical Data:No Development Reported Size:1 mg, 5 mg	$\begin{array}{c} & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & & \\ & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ &$	Xylopine is an aporphine alkaloid with cytotoxic activity on cancer cells. Xylopine induces oxidative stress, causes G2/M cell cycle arrest and apoptosis in cancer cells.         Purity:       ≥98.0%         Clinical Data:       No Development Reported         Size:       1 mg, 5 mg	O C C C C C C C C C C C C C C C C C C C
XZ739	<b>Cat. No.:</b> HY-133557	Y-27632	<b>Cat. No.:</b> HY-10071
XZ739, a <b>Cerebion</b> -dependent PROTAC <b>BCL-XL</b> ( <b>Bcl-2</b> family member) degrader with a $DC_{50}$ value of 2.5 nM in MOLT-4 cells after 16 h treatment. XZ739 also induces cell death through caspase-mediated <b>apoptosis</b> .	çoo ^{nge} rrrates	Y-27632 is an orally active, ATP-competitive inhibitor of <b>ROCK-I</b> and <b>ROCK-II</b> , with Ks of 220 and 300 nM, respectively. Y-27632 attenuates Doxorubicin-induced <b>apoptosis</b> of human cardiac stem cells.	
Purity:99.06%Clinical Data:No Development ReportedSize:5 mg, 10 mg		Purity:99.91%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 50 mg, 100 mg, 200 mg, 500 mg	
Y-27632 dihydrochloride	<b>Cat. No.:</b> HY-10583	Yatein	<b>Cat. No.:</b> HY-N1060
Y-27632 dihydrochloride is an orally active, ATP-competitive inhibitor of <b>ROCK-I</b> and <b>ROCK-II</b> , with K _i s of 220 and 300 nM, respectively. Y-27632 dihydrochloride attenuates Doxorubicin-induced <b>apoptosis</b> of human cardiac stem cells. <b>Purity:</b> 99.98% <b>Clinical Data:</b> No Development Reported <b>Size:</b> 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg	NNH	Yatein is a lignan isolated from A. chilensis, with antiproliferative activity. Yatein suppresses herpes simplex virus type 1 (HSV-1) replication by interruption the immediate-early gene expression.Purity:>98% Clinical Data:No Development Reported Size:5 mg, 10 mg, 25 mg	
YB-0158		YH-306	
(Wnt pathway inhibitor 2)	Cat. No.: HY-136541		Cat. No.: HY-120213
YB-0158 (Wnt pathway inhibitor 2) is a reverse-turn peptidomimetic and a potent <b>colorectal</b> <b>cancer stem cell (CSC)</b> targeting agent. YB-0158 disrupts Sam68-Src interactions and induces <b>apoptosis</b> in CRC cells. Anti-cancer activities.	HO HNA HNA HNA HNA HNA HNA HNA HNA HNA HNA HNA HNA	YH-306 is an antitumor agent. YH-306 suppresses colorectal tumour growth and metastasis via FAK pathway. YH-306 significantly inhibits the migration and invasion of colorectal cancer cells. YH-306 potently suppresses uninhibited proliferation and induces cell <b>apoptosis</b> .	04040-04
Clinical Data:       No Development Reported         Size:       5 mg, 10 mg, 50 mg, 100 mg		Clinical Data: No Development Reported Size: 1 mg, 5 mg	
YH239-EE	<b>Cat. No.:</b> HY-12287	ҮК-4-279	<b>Cat. No.:</b> HY-14507
YH239-EE, ethyl ester of the free carboxylic acid compound YH239, is a potent p53-MDM2 antagonizing and apoptosis-inducing agent. IC50 value: Target: MDM2/p53 YH239-EE inhibits the growth of OCI-AML-3 cells with wild type p53 by inhibiting the p53-MDM2 interaction. Purity: 99.83% Clinical Data: No Development Reported		YK 4-279 is an inhibitor of RNA Helicase A (RHA)         binding to the oncogenic transciption factor         EWS-FLI1. YK-4-279 inhibits Ewing's sarcoma family         tumor (ESFT) cell growth; YK-4-279 induces         apoptosis.         Purity:       99.61%         Clinical Data:       No Development Reported         Size       10 mM v 1 ml, 10 mg, 50 mg, 100 mg,	
Size: 10 mm × 1 mL, 10 mg, 50 mg		Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg	



Zeylenone	Cat No : HV-N2051	Zibotentan	Cat No. HV-10088
Zeylenone, a naturally occurring cyclohexene oxide, inhibits proliferation and induces <b>apoptosis</b> in cervical carcinoma cells via PI3K/AKT/mTOR and MAPK/ERK pathways.	Cat. NO. HT-N2031	Zibotentan (ZD4054) is a potent, selective and orally active <b>endothelin A</b> ( <b>ET</b> _A ) <b>receptor</b> antagonist with a K ₁ of 13 nM. Zibotentan has no inhibitory effect on ETB.	
Purity:     ≥98.0%       Clinical Data:     No Development Reported       Size:     1 mg, 5 mg		Purity:         98.19%           Clinical Data:         Phase 3           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	~ <u>N</u> ~o~
Zinc Protoporphyrin (Zn(II)-protoporphyrin IX; ZnPP; Zinc Protoporphyrin-9)	<b>Cat. No.:</b> HY-101193	ZINC69391	<b>Cat. No.:</b> HY-102078
Zinc Protoporphyrin (Zn(II)-protoporphyrin IX) is an orally active and competitive <b>heme</b> <b>oxygenase-1 (HO-1)</b> inhibitor and markedly attenuates the protective effects of Phloroglucinol (PG) against H ₂ O ₂ .		ZINC69391, a specific <b>Rac1</b> inhibitor, interferes with Rac1-GEF interaction by masking Trp56 residue on Rac1 surface. ZINC69391 interferes with the interaction of Rac1 with Dock180 and reduces Rac1-GTP levels.	
Purity:     ≥98.0%       Clinical Data:     No Development Reported       Size:     10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg	/	Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	F
Ziyuglycoside I	<b>Cat. No.</b> : HY-N0331	Ziyuglycoside II	<b>Cat. No.:</b> HY-N0332
Ziyuglycoside I isolated from S. officinalis root, has anti-wrinkle activity, and increases the expression of type I collagen. Ziyuglycoside I could be used as an active ingredient for cosmetics.	Ha George States and the second	Ziyuglycoside II is a triterpenoid saponin compound extracted from Sanguisorba officinalis L. Ziyuglycoside II induces reactive oxygen species ( <b>ROS</b> ) production and <b>apoptosis</b> . Anti-inflammation and anti-cancer effect.	HOLD HOLD HOLD HOLD HOLD HOLD HOLD HOLD
Purity:         99.47%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg		Purity:99.77%Clinical Data:No Development ReportedSize:5 mg, 10 mg	HU
ZLDI-8	<b>Cat. No.:</b> HY-123931	ZM 336372	<b>Cat. No.</b> : HY-13343
ZLDI-8 is a <b>Notch</b> activating/cleaving enzyme <b>ADAM-17</b> inhibitor and inhibits the cleavage of <b>Notch</b> protein. ZLDI-8 decreases the expression of pro-survival/anti-apoptosis and epithelial-mesenchymal transition (EMT) related proteins.		ZM 336372 is a potent inhibitor of the protein kinase c-Raf. The IC ₅₀ value is 0.07 $\mu$ M in the standard assay, which contains 0.1 mM ATP.	, ¹ O ¹ ¹ O ¹ ¹ O ₀
Purity:98.53%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg	°	Purity:     ≥96.0%       Clinical Data:     No Development Reported       Size:     10 mM × 1 mL, 5 mg, 10 mg, 50 mg	
ZM-447439	<b>Cat. No.:</b> HY-10128	ZMF-10	<b>Cat. No</b> .: HY-146786
ZM-447439 is an $aurora$ kinase inhibitor with $IC_{\rm so} s$ of 110 and 130 nM for aurora A and B, respectively.	0~;x\$ _0+0	ZMF-10 is a highly potent <b>PAK1</b> inhibitor, with $IC_{50}^{S}$ of 174 nM, 1.038 $\mu$ M and 1.372 $\mu$ M for PAK1, PAK2 and PAK3, respectively. ZMF-10 can inhibit PAK1 activity to affect PAK1-regulated <b>apoptosis</b> , ER-Stress and migration in MDA-MB-231 cells. ZMF-10 can be used for researching anticancer.	
Purity:         99.19%           Clinical Data:         No Development Reported           Size:         10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg		Purity:       >98%         Clinical Data:       No Development Reported         Size:       1 mg, 5 mg	^U N ^d H ^N
Zn(BQTC)	<b>Cat. No.:</b> HY-146287	Zoledronic Acid (Zoledronate; CGP 42446; CGP42446A; ZOL 446)	<b>Cat. No.:</b> HY-13777
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Zn(BQTC) is a highly potent <b>mitochondrial DNA</b> (mtDNA) and nuclear DNA (nDNA) inhibitor. Zn(BQTC) causes severe damage to the mtDNA and nDNA, sequentially disruptes mitochondrial and nuclear functions.		Zoledronic Acid (Zoledronate) is a third-generation bisphosphonate (BP), with potent anti-resorptive activity. Zoledronic Acid inhibits the differentiation and apoptosis of osteoclasts. Zoledronic Acid also has anti-cancer effects.	N HO POH N HO POH
Purity:>98%Clinical Data:No Development ReportedSize:1 mg, 5 mg	~~~ <u>~</u> ~~	Purity:>98%Clinical Data:LaunchedSize:1 mg, 5 mg	
Zoledronic acid monohydrate (Zoledronate monohy 42446 monohydrate; CGP42446A monohydrate;)	ydrate; CGP Cat. No.: HY-13777A	Zotatifin (eFT226)	<b>Cat. No.:</b> HY-112163
Zoledronic acid monohydrate (Zoledronate monohydrate) is a third-generation bisphosphonate (BP), with potent anti-resorptive activity. Zoledronic acid monohydrate inhibits the differentiation and apoptosis of osteoclasts.	N HO P=O N HO P=O HO OH	Zotatifin (eFT226) is a potent, selective, and well-tolerated <b>eIF4A</b> inhibitor. Zotatifin promotes eIF4A binding to specific mRNA sequences with recognition motifs in the 5'-UTRS ( $IC_{so}$ =2 nM) and interferes with the assembly of the eIF4F initiation complex.	
Purity:≥98.0%Clinical Data:LaunchedSize:50 mg, 100 mg	н ²⁰ `н	Purity:     99.58%       Clinical Data:     Phase 2       Size:     1 mg, 2 mg, 5 mg	O- OH /N-
ZРСК		ZX-29	
(SL-01)	Cat. No.: HY-100709		Cat. No.: HY-135887
ZPCK is an oral active prodrug of gemcitabine that was designed for improved oral bioavailability.	a J. H. J. J.	ZX-29 is a potent and selective ALK inhibitor with an $IC_{50}$ of 2.1 nM, 1.3 nM and 3.9 nM for ALK, ALK L1196M and ALK G1202R mutations, respectively. ZX-29 is inactive against EGFR.	
Purity: >98%   Clinical Data: No Development Reported   Size: 1 mg, 5 mg		Purity:     99.52%       Clinical Data:     No Development Reported       Size:     5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
ZZW-115	<b>Cat. No.</b> : HY-111838	ZZW-115 hydrochloride	<b>Cat. No.</b> : HY-111838A
ZZW-115 is a potent NUPR1 inhibitor, with a $K_d$ of 2.1 $\mu$ M. ZZW-115 induces tumor cell death by necroptosis and <b>apoptosis</b> . Anticancer activity.		ZZW-115 hydrochloride is a potent NUPR1 inhibitor, with a $K_d$ of 2.1 $\mu$ M. ZZW-115 hydrochloride induces tumor cell death by necroptosis and <b>apoptosis</b> . Anticancer activity.	H H Q Q A A A A A A A A A A A A A A A A
Purity:>98%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 50 mg, 100 mg	~~s~~"	Purity:98.09%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 1	00 mg
[6]-Gingerol		[8]-Shogaol	
((S)-(+)-[6]Gingerol; 6-Gingerol)	Cat. No.: HY-14615		Cat. No.: HY-N2435
-Gingerol is an active compound isolated from Ginger (Zingiber officinale Rosc), exhibits a variety of biological activities including anticancer, anti-inflammation, and anti-oxidation.	.0. С С С С С С С С С С С С С С С С С С	-Shogaol, one of the pungent phenolic compounds in ginger, exhibits anti-platelet activity ( $IC_{so}$ =5 $\mu$ M) and inhibits COX-2 ( $IC_{so}$ =17.5 $\mu$ M)Shogaol induces apoptosis in human leukemia cells.	eggelenne Hold
Purity:99.54%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 10	00 mg	Purity:99.93%Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg	

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[D-Leu-4]-OB3		α-Lipoic Acid	
[D-Leu-4]-OB3 inhibits expressions of pro-inflammatory, proliferative and metastatic genes and PD-L1 expression. [D-Leu-4]-OB3 stimulates expression of pro-apoptotic genes. Purity: >98% Clinical Data: No Development Reported	Cat. No.: HY-P3342	(Thioctic acid; (±)-α-Lipoic acid; DL-α-Lipoic acid)     α-Lipoic Acid is an antioxidant, which is an essential cofactor of mitochondrial enzyme complexes. α-Lipoic Acid inhibits NF-κB-dependent     HIV-1 LTR activation. α-Lipoic Acid induces endoplasmic reticulum (ER) stress-mediated apoptosis in hepatoma cells.     Purity:   98.03%     Clinical Data:   Launched	Cat. No.: HY-N0492
Size: 1 mg, 5 mg		Size: 10 mM × 1 mL, 500 mg	
<mark>α-Lipoic Acid-d5</mark> (Thioctic acid-d5; (±)-α-Lipoic acid-d5 DL-α-Lipoic acid-d5)	; <b>Cat. No</b> .: HY-N0492S	α-ΝΕΤΑ	<b>Cat. No.:</b> HY-138097
α-Lipoic Acid-d5 (Thioctic acid-d5) is the deuterium labeled α-Lipoic Acid. α-Lipoic Acid is an antioxidant, which is an essential cofactor of <b>mitochondrial</b> enzyme complexes. α-Lipoic Acid inhibits <b>NF-κB</b> -dependent <b>HIV-1</b> LTR activation.	D D OH	α-NETA is a potent and noncompetitive <b>choline</b> acetyltransferase (ChA) inhibitor with an IC ₅₀ of 9 μM. α-NETA is a potent ALDH1A1 (IC ₅₀ =0.04 μM) and <b>chemokine-like receptor-1 (CMKLR1)</b> antagonist.	
Purity: >98%   Clinical Data: No Development Reported   Size: 1 mg, 5 mg, 10 mg		Purity:   ≥98.0%     Clinical Data:   No Development Reported     Size:   5 mg, 10 mg, 25 mg, 50 mg, 100 mg	
α-Solanine	<b>Cat. No.:</b> HY-N6602	α-Thujone	<b>Cat. No.:</b> HY-121618
$\alpha$ -solanine, a bioactive component and one of the major steroidal glycoalkaloids in potatoes, has been observed to inhibit growth and induce <b>apoptosis</b> in cancer cells.		α-Thujone is a monoterpene isolated from Thuja occidentalis essential oil with potent anti-tumor activities. α-Thujone is a reversible modulator of the GABA type A receptor and the IC ₅₀ for α-Thujone is 21 μM in suppressing the GABA-induced currents.	0=
Purity: ≥98.0%   Clinical Data: No Development Reported   Size: 1 mg, 5 mg		Purity: ≥95.0%   Clinical Data: No Development Reported   Size: 50 mg, 100 mg	/ н
αβ-Tubulin-IN-1	<b>Cat. No.:</b> HY-144132	<mark>β-Carotene</mark> (Provitamin A; beta-Carotene)	<b>Cat. No.:</b> HY-N0411
$\alpha$ β-Tubulin-IN-1 is a potent and orally active $\alpha$ β-Tubulin inhibitor. $\alpha$ β-Tubulin-IN-1 induces cell cycle arrest at G2/M and efficient <b>apoptosis</b> . $\alpha$ β-Tubulin-IN-1 inhibits tumor cell migration and Metastasis. $\alpha$ β-Tubulin-IN-1 shows significant antitumor efficacy in a dose dependent manner.	atat Dogo	β-Carotene (Provitamin A), a carotenoid compound, is a naturally-occurring vitamin A precursor. β-Carotene is a modulator of <b>reactive oxygen</b> <b>species (ROS)</b> , with antioxidant and antiinflammatory activities.	English and the second
Purity: >98%   Clinical Data: No Development Reported   Size: 1 mg, 5 mg		Purity: ≥98.0%   Clinical Data: Launched   Size: 50 mg, 100 mg	
<mark>β-Elemene</mark> ((-)-β-Elemene; Levo-β-elemene)	<b>Cat. No.:</b> HY-107324	β-Elemonic acid	<b>Cat. No.:</b> HY-N2454
β-Elemene ((-)- $β$ -Elemene; Levo- $β$ -elemene) is isolated from natural plant Curcuma wenyujin with an antitumor activity. $β$ -Elemene can induce cell <b>apoptosis</b> .		β-Elemonic acid is a triterpene isolated from Boswellia papyrifera. β-Elemonic acid induces cell <b>apoptosis</b> , reactive oxygen species ( <b>ROS</b> ) and <b>COX-2</b> expression and inhibits <b>prolyl endopeptidase</b> . β-Elemonic acid exhibits anticancer and anti-inflammatory effects.	HOLE
Purity:     99.88%       Clinical Data:     Launched       Size:     10 mM × 1 mL, 5 mg, 10 mg, 50 mg	~~	Purity:≥99.0%Clinical Data:No Development ReportedSize:5 mg, 10 mg, 20 mg	н

β-Ionone	<b>Cat. No.:</b> HY-W015084	<mark>β-Lapachone</mark> (ARQ-501; NSC-26326)	<b>Cat. No.</b> : HY-13555
β-Ionone is effective in the induction of <b>apoptosis</b> in gastric adenocarcinoma SGC7901 cells. Anti-cancer activity.	Les i	β-Lapachone (ARQ-501;NSC-26326) is a naturally occurring O-naphthoquinone, acts as a <b>topoisomerase I</b> inhibitor, and induces apoptosis by inhibiting cell cycle progression.	e contraction of the second se
Purity: >98%   Clinical Data: No Development Reported   Size: 1 mg, 5 mg	Ň	Purity:     99.85%       Clinical Data:     Phase 2       Size:     10 mM × 1 mL, 10 mg, 50 mg, 100 mg	0
ß-Naphthoflavone		ß-NFTA	
(5,6-Benzoflavone; beta-NF)	Cat. No.: HY-114740	P	Cat. No.: HY-124957
$\beta$ -Naphthoflavone is a non-carcinogenic <b>AhR</b> agonist as a positive control for the induction of AhR transcriptional activity. $\beta$ -Naphthoflavone inhibits hydrogen peroxide-induced apoptosis.		β-NETA is a potent and noncompetitive <b>choline</b> acetyltransferase (ChA; IC ₅₀ =76 μM) and <b>cholinesterase</b> (ChE; IC ₅₀ =40 μM) inhibitor. $β$ -NETA weakly inhibits acetylcholinesterase (AChE; IC ₅₀ =1 mM).	COO ^r ,
Purity: >98%   Clinical Data: No Development Reported   Size: 1 mg, 5 mg	~~	Purity: ≥98.0%   Clinical Data: No Development Reported   Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg	
7 Stat		7 Stat trisodium	
(NSC37044)	Cat. No.: HY-123979	(NSC37044 trisodium)	Cat. No.: HY-123979A
ζ-Stat (NSC37044) is a specific and atypical PKC-ζ inhibitor, with an IC ₅₀ of 5 μM. ζ-Stat can reduce melanoma cell lines proliferation and induce apoptosis, and has antitumor activity in vitro. Purity: $\geq$ 95.0%	но он 0=5=0 но 5 о о о он	$\zeta$ -Stat trisodium (NSC37044 trisodium) is a specific and atypical PKC- $\zeta$ inhibitor, with an IC _{so} of 5 $\mu$ M. $\zeta$ -Stat trisodium can reduce melanoma cell lines proliferation and induce apoptosis, and has antitumor activity in vitro.Purity: $\geq$ 97.0%	HO QNa O=S=O O, S NaO'S O O'ONa
Clinical Data:No Development ReportedSize:10 mM × 1 mL, 5 mg, 10 mg, 50 mg		Clinical Data:No Development ReportedSize:5 mg, 10 mg, 50 mg	