

Autophagy

Autophagy is a conserved cellular degradation and recycling process in the lysosome. In mammalian cells, there are three primary types of autophagy: microautophagy, macroautophagy, and chaperone-mediated autophagy (CMA). Microphagy captures cargoes by means of invaginations or protrusions of the lysosomal membrane directly, CMA uses chaperones to identify cargo proteins and then unfolds and transfers them into the lysosomal, while macroautophagy sequesters cargo by autophagosomes-de novo synthesized of double-membrane vesicles-and subsequently transport it to the lysosome.

Macroautophagy is the best studied and it occurs at a low level constitutively and can also be further induced under stress conditions, such as nutrient or energy starvation with a salient feature of autophagy protein degradation. Stress-induced macrophagy plays an important role in protein catabolism with another key protein degradation pathway, the ubiquitin–proteasome system (UPS).

As the study progressed, autophagy gains its importance under basal, nutrient-rich conditions, and is now recognized as a critical housekeeping pathway in catabolism of diverse cellular constituents, such as protein aggregates (aggrephagy), lipid droplets (lipophagy), iron complex (Ferritinophagy) and carbohydrate. Except for macromolecules, autophagy can also target several organelles and structures, such as mitochondria (mitophagy), peroxisome (pexophagy), endoplasmic reticulum (reticulophagy or ER-phagy), ribosome (ribophagy), spermatozoon-inherited organelles following fertilization (allophagy), secretory granules within pancreatic cells (zymophagy) and intracellular pathogens (xenophagy).

Autophagy and its dysfunction are associated with a variety of human pathologies, including ageing, cancer, neurodegenerative disease, heart disease and metabolic diseases, such as diabetes. Plenty of drugs and natural products are involved in autophagy modulation through multiple signaling pathways. Small molecules that can regulate autophagy seem to have great potential to intervene such diseases in animal models or clinical courses.

Autophagy Inhibitors, Activators, Modulators, Chemicals & Inducers

(+)-JQ-1

(JQ1) Cat. No.: HY-13030

(+)-JQ-1 (JQ1) is a potent, specific, and reversible BET bromodomain inhibitor, with IC_{50} S of 77 and 33 nM for the first and second bromodomain (BRD4(1/2)). (+)-JQ-1 also activates autophagy.

S N O O

Purity: 99.90%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

(+)-Talarozole

(+)-Talarozole is a potent inhibitor of **retinoic acid** metabolism extracted from patent WO 1997049704

A1.

Rotation (+)

Cat. No.: HY-14802C

Purity: 99.28%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 25 mg, 50 mg

(-)-Epicatechin gallate

(Epicatechin gallate; ECG; (-)-Epicatechin 3-O-gallate) Cat. No.: HY-N0002

(-)-Epicatechin gallate (Epicatechin gallate) inhibits cyclooxygenase-1 (COX-1) with an IC $_{50}$ of 7.5 μM_{\odot}

HO OH OH OH

Purity: 98.57%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

(-)-Epigallocatechin

(Epigallocatechin; L-Epigallocatechin)

(-)-Epigallocatechin (Epigallocatechin) is the most abundant flavonoid in green tea, can bind to unfolded native polypeptides and prevent conversion to amyloid fibrils.

HO OH OH

Cat. No.: HY-N0225

Purity: 98.01% Clinical Data: Phase 4

Size: 10 mM × 1 mL, 5 mg, 10 mg

(-)-Epigallocatechin Gallate

(EGCG; Epigallocatechol Gallate)

(-)-Epigallocatechin Gallate is a tea flavonoid with potent antioxidant, antiinflammatory, and anticarcinogenic properties. (-)-Epigallocatechin Gallate is reported to inhibit EGFR signaling and thereby exert anticancer effects.

OH OH OH

Cat. No.: HY-13653

Purity: 99.87% Clinical Data: Phase 4

Size: $10 \text{ mM} \times 1 \text{ mL}$, 50 mg, 100 mg, 500 mg

(-)-PX20606 trans isomer

((-)-PX-102 trans isomer; (-)-PX-104)

(-)-PX20606 trans isomer is a FXR agonist with EC_{50} s of 18 and 29 nM for FXR in FRET and M1H assay, respectively.

Cat. No.: HY-100443B

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 2 mg

(-)-Rasfonin

Cat. No.: HY-121532

(-)-Rasfonin is a fungal secondary metabolite and inhibits small G proteins Ras. (-)-Rasfonin induces apoptosis, necrosis and autophagy in ACHN cells (a renal carcinoma cell line).

HO. 194

Purity: > 98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

(-)-Talarozole

(-)-Talarozole is a potent inhibitor of **retinoic** acid metabolism extracted from patent WO 1997049704

Α1

Rotation (-)

Cat. No.: HY-14802D

Purity: 98.02%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 25 mg, 50 mg

(3R,5R)-Rosuvastatin

Cat. No.: HY-17504C

(3R,5R)-Rosuvastatin is the (3R,5R)-enantiomer of Rosuvastatin. Rosuvastatin is a competitive HMG-CoA reductase inhibitor with an IC $_{50}$ of 11 nM. Rosuvastatin potently blocks human ether-a-go-go related gene (hERG) current with an IC $_{50}$ of 195 nM.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

(3R,5S)-Fluvastatin

((3R,5S)-XU 62-320 free acid)

Cat. No.: HY-14664B

(3R,5S)-Fluvastatin is the 3R,5S-isomer Fluvastatin. Fluvastatin (XU 62-320 free acid) is a first fully synthetic, competitive **HMG-CoA reductase** inhibitor with

an IC₅₀ of 8 nM.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

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(3S,5R)-Fluvastatin D6

((3S,5R)-XU 62-320 free acid D6)

Cat. No.: HY-14664DS

(3S,5R)-Fluvastatin D6 is the deuterium labeled (3S.5R)-Fluvastatin sodium. Fluvastatin is a first fully synthetic, competitive HMG-CoA reductase inhibitor with an IC₅₀ of 8 nM.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg

(E)-Daporinad

(FK866; APO866) Cat. No.: HY-50876

(E)-Daporinad (FK866) is an effective inhibitor of nicotinamide phosphoribosyltransferase (NMPRTase; Nampt) with an IC₅₀ of 0.09 nM.

Cat. No.: HY-15464

Purity: 99 94% Clinical Data: Phase 2

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

(R)-(-)-Gossypol

(AT-101; R-(-)-gossypol acetic acid)

(R)-(-)-Gossypol (AT-101) is the levorotatory isomer of a natural product Gossypol. AT-101 is determined to bind to Bcl-2, Mcl-1 and Bcl-xL proteins with K_is of 260±30 nM, 170±10 nM, and 480±40 nM, respectively.



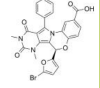
(R)-BPO-27

Size:

Cat. No.: HY-19778

(R)-BPO-27, the R enantiomer of BPO-27, is a potent, orally active and ATP-competitive CFTR inhibitor with an IC_{so} of 4 nM.

1 mg, 5 mg



99.86% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 50 mg, 100 mg Size:

(Rac)-AZD 6482

((Rac)-KIN-193) Cat. No.: HY-75124

(Rac)-AZD 6482 ((Rac)-KIN-193) is the racemate of AZD 6482. AZD 6482 is a potent and selective $p110\beta$ inhibitor with an IC_{so} of 0.69 nM.



Purity: 97.92%

No Development Reported Clinical Data:

Size 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

(3S,5R)-Rosuvastatin

(3S,5R)-Rosuvastatin is the (3S,5R)-enantiomer of Rosuvastatin, Rosuvastatin is a competitive $\overline{\text{HMG-CoA}}$ reductase inhibitor with an $\overline{\text{IC}}_{50}$ of 11 nM. Rosuvastatin potently blocks human ether-a-go-go related gene (hERG) current with an IC₅₀ of 195 nM.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

(R)-(-)-Felodipine-d5

Cat. No.: HY-132670S

Cat. No.: HY-17504D

Cat. No.: HY-15464A

(R)-(-)-Felodipine-d5 is the deuterium labeled (R)-(-)-Felodipine. (R)-(-)-Felodipine is the S enantiomer of Felodipine. Felodipine, a dihydropyridine, is a potent, vasoselective calcium channel antagonist.

Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg, 10 mg

(R)-(-)-Gossypol acetic acid (AT-101 (acetic acid);

(-)-Gossypol acetic acid; (R)-Gossypol acetic acid)

(R)-(-)-Gossypol acetic acid (AT-101 (acetic acid)) is the levorotatory isomer of a natural product Gossypol. AT-101 is determined to bind to Bcl-2, Mcl-1 and Bcl-xL proteins with K,s of 260±30 nM, 170±10 nM, and 480±40 nM, respectively.

98.02% Purity: Clinical Data: Phase 2

Size 10 mM × 1 mL, 10 mg, 50 mg

(R)-Hydroxychloroquine

((R)-HCQ) Cat. No.: HY-B1370B

(R)-Hydroxychloroquine is the enantiomer of Hydroxychloroquine. Hydroxychloroquine is a synthetic antimalarial drug which can also inhibit Toll-like receptor 7/9 (TLR7/9) signaling. Hydroxychloroquine is efficiently inhibits SARS-CoV-2 infection in vitro.

>98% Purity: Clinical Data: Launched Size 1 mg, 5 mg



Cat. No.: HY-124729A

(Rac)-BL-918 is the racemate of BL-918. BL-918 is a potent activator of UNC-51-like kinase 1 (ULK1), inducing cytoprotective autophagy for Parkinson's disease treatment.



Purity: 98.06%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

(Rac)-Efavirenz-d4

Cat. No.: HY-10572BS

(Rac)-Efavirenz-d4 ((Rac)-DMP 266-d4) is a labelled racemic Efavirenz, Efavirenz (DMP 266) is a potent inhibitor of the wild-type HIV-1 reverse transcriptase with a K_i of 2.93 nM and exhibits an IC₉₅ of 1.5 nM for the inhibition of HIV-1 replicative spread in cell culture.

Purity: >98%

Clinical Data:

Size: 1 mg, 10 mg

(Rac)-Hesperetin

(Rac)-Hesperetin is the racemate of Hesperetin. Hesperetin is a natural flavanone, and acts as a potent and broad-spectrum inhibitor against human UGT activity. Hesperetin induces apoptosis via p38 MAPK activation.



Cat. No.: HY-N0168A

98 20% Purity:

Clinical Data: No Development Reported

Size: 100 mg

(Rac)-Hesperetin-13C,d3

Cat. No.: HY-N0168AS1

(Rac)-Hesperetin-13C,d3 is the 13C- and deuterium labeled. (Rac)-Hesperetin is the racemate of Hesperetin. Hesperetin is a natural flavanone, and acts as a potent and broad-spectrum inhibitor against human UGT activity. Hesperetin induces apoptosis via p38 MAPK activation.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

(Rac)-Hesperetin-d3

Cat. No.: HY-N0168AS

(Rac)-Hesperetin-d3 is the deuterium labeled (Rac)-Hesperetin. (Rac)-Hesperetin is the racemate of Hesperetin. Hesperetin is a natural flavanone, and acts as a potent and broad-spectrum inhibitor against human **UGT** activity. Hesperetin induces apoptosis via p38 MAPK activation.

Purity:

Clinical Data: No Development Reported

1 mg, 10 mg

(Rac)-Norepinephrine-d3 (formate)

Cat. No.: HY-13715S

(Rac)-Norepinephrine-d3 (formate) is deuterium labeled Norepinephrine. Norepinephrine (Levarterenol; L-Noradrenaline) is a potent adrenergic receptor (AR) agonist. Norepinephrine activates $\alpha 1$, $\alpha 2$, $\beta 1$ receptors.

(Rac)-Sitagliptin-d4 hydrochloride

Cat. No.: HY-13749S

(Rac)-Sitagliptin-d4 hydrochloride is a labelled racemic Sitagliptin. Sitagliptin hydrochloride is a potent inhibitor of DPP4 with an IC₅₀ of 19 nM in Caco-2 cell extracts.

Purity: >98%

Clinical Data: No Development Reported

Size 1 mg, 10 mg

(S)-Hydroxychloroquine

Clinical Data: No Development Reported

Purity:

Purity:

Size: 1 mg, 5 mg

>98%

(rel)-Atorvastatin

Cat. No.: HY-B0589A

(rel)-Atorvastatin, a relative configuration of Atorvastatin. Atorvastatin is an orally active HMG-CoA reductase inhibitor, has the ability to effectively decrease blood lipids. Atorvastatin inhibits human SV-SMC proliferation and invasion with IC_{so} s of 0.39 μ M and 2.39 μ M, respectively.

((S)-HCQ)

(S)-Hydroxychloroquine ((S)-HCQ) is the enantiomer of Hydroxychloroquine. Hydroxychloroquine, a synthetic antimalarial drug, inhibits Toll-like receptor 7/9 (TLR7/9) signaling, and shows efficiently inhibits SARS-CoV-2 infection in vitro

Cat. No.: HY-B1370A

Purity: >98% Clinical Data: Launched Size 1 mg, 5 mg

Clinical Data: Launched 10 mg, 50 mg, 100 mg Size:

>98%

(S)-Sitagliptin phosphate ((S)-MK-0431 phosphate) Cat. No.: HY-13749C

(S)-Sitagliptin phosphate is the less active S-enantiomer of Sitagliptin phosphate. Sitagliptin phosphate (MK-0431 phosphate) is a potent inhibitor of DPP4 with an IC_{50} of 19 nM in Caco-2 cell extracts.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

(±)-Taxifolin

((±)-Dihydroquercetin)

(±)-Taxifolin ((±)-Dihydroquercetin) is the racemate of Taxifolin. Taxifolin exhibits important anti-tyrosinase activity. Taxifolin exhibits significant inhibitory activity against collagenase with an IC_{50} value of 193.3 μM .



Cat. No.: HY-N0136A

Purity: >98%

Clinical Data: No Development Reported

10 mg

1-Monomyristin

Cat. No.: HY-N2512

1-Monomyristin, extracted from Serenoa repens, inhibits the hydrolysis of 2-oleoylglycerol $(IC_{50}=32 \mu M)$ and fatty acid amide hydrolase (FAAH) activity (IC₅₀=18 μ M).



Purity: >98.0%

Clinical Data: No Development Reported Size: 5 mg, 10 mg, 20 mg

10058-F4

10058-F4 is a c-Myc inhibitor that prevents c-Mvc-Max dimerization and transactivation of c-Myc target gene expression.



Cat. No.: HY-12702

99 77% Purity:

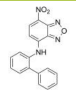
Clinical Data: No Development Reported

Size: 5 mg, 10 mg, 50 mg

10074-G5

Cat. No.: HY-100996

10074-G5 is an inhibitor of c-Myc-Max dimerization with an IC_{50} of 146 μM .



Cat. No.: HY-12033

Purity: 96.81%

2-Methoxyestradiol

(2-ME2; NSC-659853)

Clinical Data: No Development Reported

2-Methoxyestradiol (2-ME2), an orally active

endogenous metabolite of 17β-estradiol (E2), is an

potent antineoplastic activity. 2-Methoxyestradiol

apoptosis inducer and an angiogenesis inhibitor with

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg Size:

11-cis-Retinoic Acid-d5

Cat. No.: HY-14649S2

11-cis-Retinoic Acid-d5 is the deuterium labeled Retinoic acid. Retinoic acid is a metabolite of vitamin A that plays important roles in cell growth, differentiation, and organogenesis.



Purity: >98%

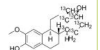
Clinical Data: No Development Reported

500 μg, 5 mg

2-Methoxyestradiol-13C6

(2-ME2-13C6; NSC-659853-13C6) Cat. No.: HY-12033S1

2-Methoxyestradiol-13C6 (2-ME2-13C6) is the 13C-labeled 2-Methoxyestradiol. 2-Methoxyestradiol (2-ME2), an orally active endogenous metabolite of 17β-estradiol (E2), is an apoptosis inducer and an angiogenesis inhibitor with potent antineoplastic activity.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Clinical Data: Phase 2 Size:

Purity:

also destablize microtubules.

99 82%

10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 500 mg

20-Deoxyingenol

20-Deoxyingenol, a diterpene, is isolated from the roots of Euphorbia kansui. 20-Deoxyingenol can promote autophagy and lysosomal biogenesis by promoting the nuclear translocation of transcription factor EB (TFEB) in vitro.



Cat. No.: HY-N0866

Purity: 99.84%

Clinical Data: No Development Reported 10 mM \times 1 mL, 5 mg, 10 mg Size

2-Methoxyestradiol-d5 (2-ME2-d5; NSC-659853-d5)

2-Methoxyestradiol-d5 is the deuterium labeled 2-Hydroxyestradiol. 2-Methoxyestradiol (2-ME2), an orally active endogenous metabolite of 17β-estradiol (E2), is an apoptosis inducer and an angiogenesis inhibitor with potent antineoplastic

activity.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

3'-Hydroxypterostilbene

Cat. No.: HY-N6002

Cat. No.: HY-12033S2

3'-Hydroxypterostilbene, a natural pterostilbene analogue, effectively inhibits the growth of human colon cancer cells (IC_{so}s of 9.0, 40.2, and 70.9 μ M for COLO 205, HCT-116, and HT-29 cells, respectively) by inducing apoptosis and autophagy.

Purity: 99.46%

Clinical Data: No Development Reported $10 \text{ mM} \times 1 \text{ mL}, 5 \text{ mg}, 10 \text{ mg}, 20 \text{ mg}$ Size

3,3'-Diindolylmethane

(DIM; Arundine; HB 236)

3,3'-Diindolylmethane is a strong, pure androgen receptor (AR) antagonist.



Cat. No.: HY-15758

98.78% Clinical Data: Phase 4

10 mM × 1 mL, 100 mg, 200 mg, 500 mg

3-Bromopyruvic acid

(Bromopyruvic acid; Hexokinase II Inhibitor II, 3-BP)

Cat. No.: HY-19992

3-Bromopyruvate (Bromopyruvic acid) is an analogue of pyruvate and a potent hexokinase (HK)-II inhibitor with high tumor selectivity. 3-Bromopyruvate inhibits cell growth and induces apoptosis through interfering with glycolysis.

Purity: 98 00%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g, 10 g, 25 g

3-Methyladenine (3-MA)

3-Methyladenine (3-MA) is a PI3K inhibitor. 3-Methyladenine is a widely used inhibitor of autophagy via its inhibitory effect on class III



Cat. No.: HY-19824

Cat. No.: HY-19312

99 83% Purity:

Clinical Data: No Development Reported

Size: 50 mg, 100 mg, 200 mg, 500 mg

3BDO

Cat. No.: HY-U00434

3BDO is a new mTOR activator which can also inhibit autophagy.

Purity: 99 91%

Clinical Data: No Development Reported 10 mM × 1 mL, 5 mg, 10 mg, 25 mg

3PO

3PO is a novel small-molecule inhibitor of the PFKFB3 isozyme, 3PO markedly attenuates the proliferation of several human malignant hematopoietic and adenocarcinoma cell lines (IC50, 1.4-24 μM) IC50 value Target: PFKFB3 isozyme in

vitro: 3PO inhibits recombinant PFKFB3...

Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

4,4'-Dimethoxychalcone

Cat. No.: HY-136064

4,4'-Dimethoxychalcone acts as a natural autophagy inducer with anti-ageing properties.

Purity: ≥98.0%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 25 mg, 50 mg, 100 mg

4-Hydroxylonchocarpin

4-Hydroxylonchocarpin is a chalcone compound from an extract of Psoralea corylifolia.

4-Hydroxylonchocarpin increases phosphorylation of p38 MAPK, JNK and ERK.

Cat. No.: HY-N2208

Purity: 92.14%

Clinical Data: No Development Reported

Size 5 mg, 10 mg

4-Hydroxytolbutamide

(Hydroxytolbutamide) Cat. No.: HY-100641

4-Hydroxytolbutamide (Hydroxytolbutamide) is a metabolite of Tolbutamide. 4-Hydroxytolbutamide is metabolized by CYP2C8 and CYP2C9. Tolbutamide is a first generation potassium channel blocker and a sulfonylurea oral antidiabetic.

>98% Purity:

Clinical Data: No Development Reported

Size 5 mg, 10 mg

4-Hydroxytolbutamide-d9

(Hydroxytolbutamide-d9)

4-Hydroxytolbutamide-d9 (Hydroxytolbutamide-d9) is the deuterium labeled 4-Hydroxytolbutamide.

4-Hydroxytolbutamide (Hydroxytolbutamide) is a metabolite of Tolbutamide. 4-Hydroxytolbutamide is metabolized by CYP2C8 and CYP2C9.

Cat. No.: HY-100641S

Purity: >98%

Clinical Data: No Development Reported

Size 1 mg, 10 mg

4E2RCat

Cat. No.: HY-100733

4E2RCat is an inhibitor of eIF4E-eIF4G interaction with an IC_{so} of 13.5 μ M.



≥98.0%

Clinical Data: No Development Reported

10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 50 mg, 100 mg

4E1RCat

Cat. No.: HY-14427

4E1RCat is an inhibitor of cap-dependent translation, and inhibits eIF4E:eIF4GI interaction, with an IC_{so} an of 4 μM.



Purity: 99.10%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

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4EGI-1

Cat. No.: HY-19831

4EGI-1 is an inhibitor of eIF4E/eIF4G interaction, with a K_a of 25 μM against eIF4E binding.

98 83% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

5-Aminolevulinic acid-15N hydrochloride (5-ALA-15N

hydrochloride; ...) Cat. No.: HY-N0305S

5-Aminolevulinic acid-15N (5-ALA-15N) hydrochloride is the 15N-labeled 5-Aminolevulinic acid (hydrochloride). 5-Aminolevulinic acid hydrochloride (5-ALA hydrochloride) is an intermediate in heme biosynthesis in the body and the universal precursor of tetrapyrroles.

Purity:

Size: 1 mg, 5 mg

Clinical Data: No Development Reported

5-Aminolevulinic acid hydrochloride (5-ALA hydrochloride; Cat. No.: HY-N0305

δ-Aminolevulinic acid hydrochloride; ...)

5-Aminolevulinic acid hydrochloride (5-ALA hydrochloride) is an intermediate in heme biosynthesis in the body and the universal precursor of tetrapyrroles.



Purity: >97.0% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 1 g, 5 g, 10 g

5-Aminolevulinic acid-d2 hydrochloride (5-ALA-d2

hydrochloride; ...) Cat. No.: HY-N0305S1

5-Aminolevulinic acid-d2 (hydrochloride) is deuterium labeled 5-Aminolevulinic acid (hydrochloride).



Purity: >98% Clinical Data

1 mg, 5 mg

5-Azacytidine

(Azacitidine; 5-AzaC; Ladakamycin) Cat. No.: HY-10586

5-Azacytidine (Azacitidine; 5-AzaC; Ladakamycin) is a nucleoside analogue of cytidine that specifically inhibits DNA methylation.



Purity: 99 40% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 200 mg, 500 mg

6-Mercaptopurine

(Mercaptopurine; 6-MP)

6-Mercaptopurine is a purine analogue which acts as an antagonist of the endogenous purines and has been widely used as antileukemic agent and immunosuppressive drug.



Cat. No.: HY-13677

99.16% Purity: Clinical Data: Launched

Size 10 mM × 1 mL, 50 mg, 100 mg, 500 mg

6-Mercaptopurine-13C2,15N

(Mercaptopurine-13C2,15N; 6-MP-13C2,15N) Cat. No.: HY-13677S1

6-Mercaptopurine-13C2,15N (Mercaptopurine-13C2,15N) is the 13C- and 15N-labeled 6-Mercaptopurine. 6-Mercaptopurine is a purine analogue which acts as an antagonist of the endogenous purines and has been widely used as antileukemic agent and immunosuppressive drug.



Purity:

Clinical Data: No Development Reported

Size 1 mg, 5 mg

6-Mercaptopurine-d2

(Mercaptopurine-d2; 6-MP-d2)

6-Mercaptopurine-d2 (Mercaptopurine-d2) is the deuterium labeled 6-Mercaptopurine. 6-Mercaptopurine is a purine analogue which acts as an antagonist of the endogenous purines and has been widely used as antileukemic agent and immunosuppressive drug.



Cat. No.: HY-13677S

Purity: >98%

Clinical Data: No Development Reported Size: 2.5 mg, 1 mg, 5 mg, 10 mg

6-Thioguanine

(Thioguanine; 2-Amino-6-purinethiol) Cat. No.: HY-13765

6-Thioguanine (Thioguanine; 2-Amino-6-purinethiol) is an anti-leukemia and immunosuppressant agent, acts as an inhibitor of SARS and MERS coronavirus papain-like proteases (PLpros) and also potently inhibits USP2 activity, with IC_{50} s of 25 μM and 40 μM for Plpros and recombinant human...



Purity: ≥99.0% Clinical Data: Launched

Size 10 mM × 1 mL, 100 mg, 500 mg

740 Y-P

(740YPDGFR; PDGFR 740Y-P)

740 Y-P (740YPDGFR; PDGFR 740Y-P) is a potent and cell-permeable PI3K activator. 740 Y-P readily binds GST fusion proteins containing both the Nand C- terminal SH2 domains of p85 but fails to bind GST alone.

Cat. No.: HY-P0175

Purity: 99.67%

Clinical Data: No Development Reported

1 mg, 5 mg

740 Y-P TFA

(740YPDGFR TFA; PDGFR 740Y-P TFA)

740 Y-P TFA is a potent and cell-permeable PI3K activator, 740 Y-P TFA readily binds GST fusion proteins containing both the N- and C- terminal SH2 domains of p85 but fails to bind GST alone.

Cat. No.: HY-P0175A

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

8-Aminoadenosine

(8-NH2-Ado)

8-Aminoadenosine (8-NH2-Ado), a RNA-directed nucleoside analogue, reduces cellular ATP levels and inhibits mRNA synthesis. 8-Aminoadenosine blocks Akt/mTOR signaling and induces autophagy and apoptosis in a p53-independent manner. 8-Aminoadenosine has antitumor activity.

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg



Cat. No.: HY-125927

9-cis,13-cis-Retinoic acid-d5

Cat. No.: HY-15127S2

9-cis,13-cis-Retinoic acid-d5 is the deuterium labeled Isotretinoin, Isotretinoin (13-cis-Retinoic acid) is a medication used for the treatment of severe acne.

Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg

9-ING-41

Cat. No.: HY-113914

9-ING-41 is a maleimide-based ATP-competitive and selective glycogen synthase kinase-3ß (GSK-3ß) inhibitor with an IC_{50} of 0.71 μ M. 9-ING-41 significantly leads to cell cycle arrest, autophagy and apoptosis in cancer cells.

Purity: 99.32% Clinical Data: Phase 2

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg



A-484954

Cat. No.: HY-110096

A-484954 is a highly selective eukaryotic elongationfactor-2 (eEF2) inhibitor, with an IC₅₀ of 280 nM.

Purity: 98.10%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg Size:

A-867744

Cat. No.: HY-12149

A-867744 is a highly potent and selective type ${\rm II}$ positive allosteric modulator (PAM) of the alpha7 nicotinic acetylcholine receptors (nAChR) with an EC_{50} of 1.0 μ M.



Cat. No.: HY-50907

Cat. No.: HY-U00141

99.92% Purity:

Clinical Data: No Development Reported

Size 10 mM × 1 mL, 10 mg, 50 mg, 100 mg

ABT 737-d8

Cat. No.: HY-50907S

ABT 737-d8 is the deuterium labeled ABT-737. ABT-737, a BH3 mimetic, is a potent Bcl-2, Bcl-x, and Bcl-w inhibitor with EC₅₀s of 30.3 nM, 78.7 nM, and 197.8 nM, respectively.



Purity: >98%

Clinical Data: No Development Reported

Size 1 mg, 10 mg ABT-737

Purity:

ABT-737, a BH3 mimetic, is a potent Bcl-2, Bcl-x_L and Bcl-w inhibitor with EC₅₀s 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



Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

ABT-751

(E7010)Cat. No.: HY-13270

ABT-751(E 7010) is a novel bioavailable tubulin-binding and antimitotic sulfonamide agent with IC50 of about 1.5 and 3.4 μM in neuroblastoma and non-neuroblastoma cell lines, respectively.



Purity: 99.93% Clinical Data: Phase 2

10 mM × 1 mL, 10 mg, 50 mg, 100 mg Size:

ABTL-0812

(α-Hydroxylinoleic acid)

ABTL-0812 (α-Hydroxylinoleic acid) induces endoplasmic reticulum (ER) stress-mediated autophagy. ABTL-0812 is a first-in-class small

molecule with anti-cancer activity.

98.06% Clinical Data: Phase 2

5 mg, 10 mg, 50 mg, 100 mg

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AC-55649

AC-55649 is a potent, highly isoform-selective agonist of human RARβ2 receptor, with a pEC_{so}

of 6.9.

....ог^јон

Cat. No.: HY-108526

Purity: 99.93%

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 5 mq, 10 mq, 25 mq

AC-73

AC-73 is a first specific, orally active inhibitor of cluster of differentiation 147 (CD147), which specifically disrupts CD147 dimerization, thereby mainly suppressing the CD147/ERK1/2/STAT3/MMP-2 pathways. AC-73 inhibits the motility and invasion of hepatocellular carcinoma cells.

HO OF HOO

Cat. No.: HY-122214

Purity: 99.72%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Acacetin

(5,7-Dihydroxy-4'-methoxyflavone)

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 PI3Kγ. Acacetin causes cell cycle arrest and induces apoptosis and autophagy in cancer cells.

Cat. No.: HY-N0451

Purity: 99.84%

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 5 mq, 10 mq, 25 mq

Aceglutamide

(α-N-Acetyl-L-glutamine; N2-Acetylglutamine)

Aceglutamide (α -N-Acetyl-L-glutamine) is a psychostimulant and nootropic, used to improve memory and concentration.



Cat. No.: HY-B1065

Purity: ≥98.0% Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 1 g

Acetazolamide

Cat. No.: HY-B0782

Acetazolamide is a **carbonic anhydrase** (CA) IX inhibitor with an $\rm IC_{50}$ of 30 nM for hCA IX. Diuretic effects.

Purity: 99.97% Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g

Acetazolamide-13C2,d3

Cat. No.: HY-B0782S1

Acetazolamide-13C2,d3 is the 13C- and deuterium labeled. Acetazolamide is a carbonic anhydrase (CA) IX inhibitor with an IC50 of 30 nM for hCA IX. Diuretic effects.



Purity: >98% Clinical Data:

Size: 1 mg, 5 mg

Acitretin

(Ro 10-1670) Cat. No.: HY-B0107

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.

Purity: 99.79%
Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 200 mg, 500 mg

Acitretin sodium

(Ro 10-1670 sodium) Cat. No.: HY-B0107A

Acitretin (Ro 10-1670) sodium is a second-generation, systemic retinoid that has been used in the treatment of psoriasis. Acitretin sodium also can be used for the research of Alzheimer's disease.

Purity: >98%
Clinical Data: Launched
Size: 1 mg, 5 mg

Acitretin-d3

(Ro 10-1670-d3) Cat. No.: HY-B0107S

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.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Actein

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.



Cat. No.: HY-N6872

Purity: 98.58%

Clinical Data: No Development Reported

Size: 5 mg

Acumapimod

(BCT197) Cat. No.: HY-16715

Acumapimod (BCT197) is an orally active p38 MAP kinase inhibitor, with an IC_{so} of less than $1 \mu M$ for p38 α .

99 63% Purity: Clinical Data: Phase 2

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

(CD271)

Adapalene

Cat. No.: HY-B0091

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 RARB, RARy, RARα, respectively.



Purity: >98.0% Clinical Data: Launched

Size: 10 mM × 1 mL, 50 mg, 100 mg, 500 mg

Adapalene sodium salt

(CD 271 sodium salt) Cat. No.: HY-B0091A

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 AC₅₀s of 2.3 nM, 9.3 nM, and 22 nM for RARβ, RARγ, RARα, respectively.



Purity: >98% Clinical Data: Launched 1 mg, 5 mg

Adapalene-d3

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 AC₅₀s of 2.3 nM, 9.3 nM, and 22 nM for RARB, RARY, RARα, respectively.

Purity: >98%

Clinical Data: No Development Reported

1 mg, 10 mg Size:



Cat. No.: HY-B0091S

Adapalene-d6 Methyl Ester

Cat. No.: HY-B0091S1

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 RARβ, RARγ, RARα, respectively.

Purity: >98%

Clinical Data: No Development Reported

Size: 10 mg, 100 mg

Adenosine

(Adenine riboside; D-Adenosine)

Adenosine (Adenine riboside), a ubiquitous endogenous autacoid, acts through the enrollment of four G protein-coupled receptors: A1, A2A, A2B,



Cat. No.: HY-B0228

Size 10 mM \times 1 mL, 500 mg, 1 g, 5 g

99 92% Purity: Clinical Data: Launched

Adenosine 5'-diphosphoribose sodium

(ADP ribose sodium) Cat. No.: HY-100973A

Adenosine 5'-diphosphoribose sodium (ADP ribose sodium) is a nicotinamide adenine nucleotide (NAD+) metabolite. Adenosine 5'-diphosphoribose sodium is the most potent and primary intracellular Ca2+-permeable cation TRPM2 channel activator.



Purity: 99.03%

Clinical Data: No Development Reported

Size 10 ma

Adenosine-d1

(Adenine riboside-d1: D-Adenosine-d1) Cat. No.: HY-B0228S

Adenosine-d1 (Adenine riboside-d1) is the deuterium labeled Adenosine. Adenosine (Adenine riboside), a ubiquitous endogenous autacoid, acts through the enrollment of four G protein-coupled receptors: A1, A2A, A2B, and A3.



Purity: >98%

Clinical Data: No Development Reported

Size 1 mg, 5 mg

Adezmapimod

(SB 203580; RWJ 64809) Cat. No.: HY-10256

Adezmapimod (SB 203580) is a selective and ATP-competitive p38 MAPK inhibitor with IC_{so}s of 50 nM and 500 nM for SAPK2a/p38 and SAPK2b/p38β2, respectively. Adezmapimod inhibits LCK, GSK3 β and PKB α with IC₅₀s of 100-500-fold higher than that for SAPK2a/p38.



Purity: 99.92%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg

Adezmapimod hydrochloride

(SB 203580 hydrochloride; RWJ 64809 hydrochloride)

Adezmapimod (SB 203580) hydrochloride is a selective and ATP-competitive p38 MAPK inhibitor with IC_{so}s of 50 nM and 500 nM for SAPK2a/p38 and SAPK2b/p38β2, respectively.



Cat. No.: HY-10256A

Purity: 99.71%

Clinical Data: No Development Reported

10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg

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Afatinib

(BIBW 2992) Cat. No.: HY-10261

Afatinib (BIBW 2992) is an irreversible EGFR family inhibitor with $\rm IC_{s0}s$ of 0.5 nM, 0.4 nM, 10 nM and 14 nM for EGFR**, EGFRL858R, EGFRL858R/T990M and HER2, respectively.



Purity: 99.93% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg

Afatinib D6

(BIBW 2992 D6) Cat. No.: HY-10261S

Afatinib D6 (BIBW 2992 D6) is deuterium labeled Afatinib. Afatinib (BIBW 2992) is an irreversible EGFR family inhibitor.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Afatinib dimaleate

(BIBW 2992MA2) Cat. No.: HY-10261A

Afatinib dimaleate is an irreversible EGFR family inhibitor with IC_{50} s of 0.5 nM, 0.4 nM, 10 nM and 14 nM for EGFR*, FSPECtively.



Purity: 99.61%
Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg

Afatinib impurity 11

Cat. No.: HY-133780

Afatinib impurity 11 is an impurity of Afatinib. Afatinib is an irreversible EGFR family inhibitor with IC $_{50}$ S of 0.5 nM, 0.4 nM, 10 nM and 14 nM for EGFR $^{\text{tx}}$, EGFR $^{\text{L858R}}$, EGFR $^{\text{L858R}}$ /T790M and HER2, respectively.



Purity: 99.10%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Afatinib-d4

(BIBW 2992-d4) Cat. No.: HY-10261S1

Afatinib-d4 (BIBW 2992-d4) is the deuterium labeled Afatinib. Afatinib (BIBW 2992) is an irreversible EGFR family inhibitor with $\rm IC_{so}^s$ of 0.5 nM, 0.4 nM, 10 nM and 14 nM for EGFR^{wt}, EGFR^{LSSR}, EGFR^{LSSR/T790M} and HER2, respectively.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Afatinib-d6 dimaleate

(BIBW 2992MA2-d6) Cat. No.: HY-10261AS

Afatinib-d6 dimaleate (BIBW 2992MA2-d6) is the deuterium labeled Afatinib dimaleate. Afatinib dimaleate is an irreversible EGFR family inhibitor with $\rm IC_{50}S$ of 0.5 nM, 0.4 nM, 10 nM and 14 nM for EGFR^{WF}, EGFR^{L858R}, EGFR^{L858R}/, EGFR^{L858R}/, respectively.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Afzelin

(Kaempferol-3-O-rhamnoside) Cat. No.: HY-N1441

Afzelin (Kaempferol-3-O-rhamnoside) is is a flavonol glycoside found in Houttuynia cordata Thunberg and is widely used in the preparation of antibacterial and antipyretic agents, detoxicants and for the treatment of inflammation.



Purity: 99.62%

Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg, 25 mg

AG490

(Tyrphostin AG490; Tyrphostin B42) Cat. No.: HY-12000

AG490 (Tyrphostin AG490) is a tyrosine kinase inhibitor that inhibits EGFR, Stat-3 and IAK2/3



Purity: 99.92%

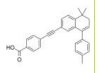
Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg

AGN 193109

Cat. No.: HY-U00449

AGN 193109 is a retinoid analog, and acts as a specific and highly effective antagonist of retinoic acid receptors (RARs), with $K_a s$ of 2 nM, 2 nM, and 3 nM for RAR α , RAR β , and RAR γ , respectively.



Purity: 99.31%

Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg

AGN 193109-d7

Cat. No.: HY-U00449S

AGN 193109-d7 is the deuterium labeled AGN 193109. AGN 193109 is a retinoid analog, and acts as a specific and highly effective antagonist of retinoic acid receptors (RARs), with $K_{\text{d}}s$ of 2 nM, 2 nM, and 3 nM for RAR α , RAR β , and RAR γ , respectively.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 10 mg

AGN 194078

Cat. No.: HY-100273

AGN 194078 is a selective RARα agonist with a K_d and EC_{so} of 3 and 112 nM, respectively.

Purity: 98 00%

Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg, 25 mg, 50 mg

AGN 194310

(VTP-194310)

AGN 194310 (VTP-194310) is a high affinity, potent and selective retinioic acid receptors (RARs) pan-antagonist with K_d values of 3 nM, 2 nM, 5 nM for RARα, RARβ, RARγ, respectively.



Cat. No.: HY-16681

98.01% Purity:

Clinical Data: No Development Reported

1 mg, 5 mg, 10 mg, 25 mg, 50 mg

AGN 196996

Cat. No.: HY-16682

AGN 196996 is a potent and selective RARα antagonist with Ki value of 2 nM; little binding affinity for RARβ(Ki=1087 nM) and RARγ(Ki=8523 nM).

Purity: 99 37%

Clinical Data: No Development Reported

1 mg, 5 mg

AGN 205327

AGN 205327 is a potent synthetic RARs agonist with EC50 of 3766/734/32 nM for RAR $\alpha/\beta/\gamma$ respectively; no inhibition on RXR. IC50 value: 3766/734/32 nM

for RAR $\alpha/\beta/\gamma$ Target: RAR agonist.



Cat. No.: HY-16685

Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg

AGN 205728

Cat. No.: HY-16683

AGN 205728 is a potent and selective RARy antagonist with Ki/IC95 values of 3 nM/ 0.6 nM; no inhibiton on RARα and RARβ. IC50 value: 3 nM/ 0.6 nM(Ki/IC95) Target: RARy antagonist More information can be found in the following patent, Compound 7a.



Purity: 96.66%

Clinical Data: No Development Reported

Size: 1 ma, 5 ma

AGN-195183

(IRX-5183; VTP-195183; NRX-195183)

AGN-195183 (IRX-5183) is a potent and selective agonist of RAR α (K_d =3 nM) with improved binding selectivity relative to AGN 193836. AGN-195183 has no activity on RARβ/γ.



Cat. No.: HY-16684

99.76% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg Size

AICAR

(Acadesine; AICA Riboside)

AICAR (Acadesine) is an adenosine analog and a AMPK activator. AICAR regulates the glucose and lipid metabolism, and inhibits proinflammatory cytokines and iNOS production. AICAR is also an autophagy, YAP and mitophagy inhibitor.



Cat. No.: HY-13417

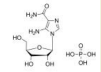
99.92% Purity: Clinical Data: Phase 3

Size: 50 mg, 100 mg, 200 mg, 500 mg

AICAR phosphate

(Acadesine phosphate; AICA Riboside phosphate)

AICAR phosphate (Acadesine phosphate) is an adenosine analog and a AMPK activator. AICAR phosphate regulates the glucose and lipid metabolism, and inhibits proinflammatory cytokines and iNOS production. AICAR phosphate is also an autophagy, YAP and mitophagy inhibitor.



Cat. No.: HY-13417A

99.49% Purity: Clinical Data: Phase 3

10 mM × 1 mL, 50 mg, 100 mg, 200 mg, 500 mg

AL 8697

Cat. No.: HY-108645

AL 8697 is a specific and orally active $p38\alpha$ \mbox{MAPK} inhibitor with an $\mbox{IC}_{\mbox{\scriptsize 50}}$ of 6 nM. AL 8697 displays 14-fold greater inhibition of p38α compared to p38 β (IC₅₀=82 nM), and 300-fold selectivity for p38α over a panel of 91 kinases. Anti-inflammatory activity.



99.49% Purity:

Clinical Data: No Development Reported $10 \text{ mM} \times 1 \text{ mL}, 5 \text{ mg}$ Size

Alginic acid

Alginic acid is a natural polysaccharide, which has been widely concerned and applied due to its excellent water solubility, film formation, biodegradability and biocompatibility.



Cat. No.: HY-W127758

>98% Purity:

Clinical Data: No Development Reported

1 mg, 5 mg

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Alisertib

(MLN 8237) Cat. No.: HY-10971

Alisertib (MLN 8237) is an orally active and selective Aurora A kinase inhibitor (IC_{so}=1.2 nM), which binds to Aurora A kinase resulting in mitotic spindle abnormalities, mitotic accumulation.



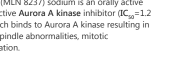
Purity: 99 84% Clinical Data: Phase 3

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

Alisertib sodium

(MLN 8237 sodium) Cat. No.: HY-10971A

Alisertib (MLN 8237) sodium is an orally active and selective Aurora A kinase inhibitor (IC₅₀=1.2 nM), which binds to Aurora A kinase resulting in mitotic spindle abnormalities, mitotic accumulation.



>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg



Aliskiren

(CGP 60536; CGP60536B; SPP 100) Cat. No.: HY-12176

Aliskiren(CGP 60536) is a direct renin inhibitor with IC50 of 15 nM

Purity: 99 16% Clinical Data: Launched

10 mM × 1 mL, 5 mg, 10 mg, 50 mg

Aliskiren hemifumarate (CGP 60536 hemifumarate; CGP60536B

hemifumarate; SPP 100 hemifumarate) Cat. No.: HY-12177

Aliskiren hemifumarate(CGP 60536 hemifumarate) is a direct renin inhibitor with IC50 of 1.5 nM



Purity: 98 98% Clinical Data: Launched

10 mM × 1 mL, 10 mg, 50 mg, 100 mg

Aliskiren-d6 hemifumarate (CGP 60536 D6 hemifumarate;

CGP60536B D6 hemifumarate; SPP 100 D6 hemifumarate) Cat. No.: HY-12177S

Aliskiren D6 hemifumarate (CGP 60536 D6 hemifumarate) is a deuterium labeled Aliskiren hemifumarate. Aliskiren hemifumarate is a direct and orally active renin inhibitor with an IC_{so} of 1.5 nM.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Alisol A

(Alisol-A)

Alisol A is a natural product.



Cat. No.: HY-N0853

99.22% Purity:

Clinical Data: No Development Reported

Size: 5 mg, 10 mg

ALLO-1

Cat. No.: HY-121546

ALLO-1, an autophagy receptor, is essential for autophagosome formation around paternal organelles and directly binds to the worm LC3 homologue LGG-1 through its LC3-interacting region (LIR) motif.



Purity: >98%

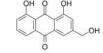
Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Aloe emodin

(Rhabarberone; 3-Hydroxymethylchrysazine)

Aloe emodin is a hydroxyanthraquinone present in Aloe vera leaves, has a specific in vitro and in vivo antitumor activity. IC50 value: Target: in vitro: aloe-emodin treatment led to the dissociation of heat shock protein 90 (HSP90) and ER α and increased ER α ubiquitination.



Cat. No.: HY-N0189

Purity: 98.32%

Clinical Data: No Development Reported

100 mg, 500 mg Size

Aloperine

Cat. No.: HY-13516

Aloperine is an alkaloid in sophora plants such as Sophora alopecuroides L, which has shown anti-cancer, anti-inflammatory and anti-virus properties. Aloperine is widely used to treat patients with allergic contact dermatitis eczema and other skin inflammation in China.



Purity: ≥98.0%

Clinical Data: No Development Reported Size: 10 mM × 1 mL, 50 mg

AM580

(CD336; NSC608001; Ro 40-6055)

AM580 is a selective RAR α agonist with IC₅₀ and EC₅₀ of 8 nM and 0.36 nM, respectively.

Cat. No.: HY-10475

99.61%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Ambroxol

(NA-872) Cat. No.: HY-B1039

Ambroxol (NA-872), an active metabolite of the prodrug Bromhexine, has potent expectorant effects. Ambroxol is a glucocerebrosidase (GCase) chaperone and increases glucocerebrosidase activity.

Purity: 99.83% Clinical Data: Launched

Size: $10 \text{ mM} \times 1 \text{ mL}, 100 \text{ mg}$

Ambroxol hydrochloride

(NA-872 hydrochloride)

Ambroxol hydrochloride (NA-872 hydrochloride), an active metabolite of the prodrug Bromhexine, has potent expectorant effects. Ambroxol hydrochloride is a glucocerebrosidase (GCase) chaperone and increases glucocerebrosidase activity.



Cat. No.: HY-B1039A

≥98.0% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg

Ambroxol-d5 hydrochloride

(NA-872-d5 hydrochloride) Cat. No.: HY-B1039AS

Ambroxol-d5 (NA-872-d5) hydrochloride is the deuterium labeled Ambroxol hydrochloride. Ambroxol hydrochloride (NA-872 hydrochloride), an active metabolite of the prodrug Bromhexine, has potent expectorant effects.

Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg

AMG PERK 44

AMG PERK 44 is an orally active and highly selective PERK inhibitor with an IC₅₀ of 6 nM. AMG PERK 44 has 1000-fold and 160-fold selectivity over GCN2 (IC_{50} =7300 nM) and B-Raf (IC_{50} >1000 nM), respectively. AMG PERK 44 induces autophagy.

Cat. No.: HY-12661A

Purity: 99.17%

Clinical Data: No Development Reported 5 mg, 10 mg, 50 mg

Amiodarone

Cat. No.: HY-14187

Amiodarone is an antiarrhythmic drug for inhibition of ATP-sensitive potassium channel with an IC $_{so}$ of 19.1 μ M.



Purity: >98% Clinical Data: Launched Size: 1 mg, 5 mg

Amiodarone hydrochloride

Cat. No.: HY-14188

Amiodarone hydrochloride, a benzofuran-based Class III antiarrhythmic agent, inhibits WT outwardIhERG tails with an IC_{so} of 45 nM.



99.96% Purity: Clinical Data: Launched

Size 10 mM \times 1 mL, 500 mg, 1 g, 5 g

Amiodarone-d10 hydrochloride

Cat. No.: HY-14187S

Amiodarone-d10 hydrochloride is the deuterium labeled Amiodarone. Amiodarone hydrochloride is an antiarrhythmic drug for inhibition of ATP-sensitive potassium channel with an IC_{so} of 19.1 μΜ.



>98% Purity:

Clinical Data: No Development Reported 1 mg, 5 mg, 10 mg, 25 mg, 50 mg Size:

Amiodarone-d4 hydrochloride

Amiodarone-d4 hydrochloride is the deuterium labeled Amiodarone hydrochloride. Amiodarone

hydrochloride, a benzofuran-based Class III antiarrhythmic agent, inhibits WT outwardIhERG tails with an IC_{50} of 45 nM.

Purity: >98%

Clinical Data: No Development Reported

Size 5 ma



Cat. No.: HY-14188S

Ammonium chloride

>98%

250 mg

Clinical Data: Launched

Purity:

Size:

Cat. No.: HY-Y1269

pH value regulation, can cause intracellular alkalization and metabolic acidosis thus effecting enzymatic activity and influencing the process of biological system. Ammonium chloride is an autophagy inhibitor.



Ammonium chloride, as a heteropolar compound with

Amsacrine

(m-AMSA; acridinyl anisidide)

Amsacrine (m-AMSA; acridinyl anisidide) is an inhibitor of topoisomerase II, and acts as an antineoplastic agent which can intercalates into the DNA of tumor cells.



Cat. No.: HY-13551

Purity: 99.98% Clinical Data: Launched

10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 500 mg

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Amsacrine hydrochloride

(m-AMSA hydrochloride; acridinyl anisidide hydrochloride) Cat. No.: HY-13551A

Amsacrine hydrochloride (m-AMSA hydrochloride; acridinyl anisidide hydrochloride) is an inhibitor of **topoisomerase II**, and acts as an antineoplastic agent which can intercalates into the DNA of tumor cells.

Purity: 98.98% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 500 mg

$\label{lem:condition} \textbf{Ancitabine hydrochloride} \hspace{0.2cm} \textbf{(Cyclocytidine hydrochloride;}$

Cyclo-CMP hydrochloride; Cyclo-C)

Cat. No.: HY-N0093

Ancitabine (hydrochloride) is an important antileukemia drugs.

Purity: 98.97% Clinical Data: Launched

Size: 10 mM × 1 mL, 200 mg, 1 g

Andrographolide

(Andrographis) Cat. No.: HY-N0191

Andrographolide is a NF- κ B inhibitor, which inhibits NF- κ B activation through covalent modification of a cysteine residue on p50 in endothelial cells without affecting I κ B α degradation or p50/p65 nuclear translocation. Andrographolide has antiviral effects.



Purity: 98.57%
Clinical Data: Launched
Size: 100 mg, 500 mg

Apigenin (4',5,7-Trihydroxyflavone; Apigenol; C.I. Natural

Yellow 1)

w 1) Cat. No.: HY-N1201

Apigenin (4',5,7-Trihydroxyflavone) is a competitive CYP2C9 inhibitor with a K_i of 2 μM .



Purity: 99.22%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg, 500 mg

Apocynin

(Acetovanillone) Cat. No.: HY-N0088

Apocynin is a selective NADPH-oxidase inhibitor with an IC_{50} of 10 μ M.

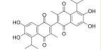
Purity: 99.95% Clinical Data: Phase 1

Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g

Apogossypolone

(ApoG2) Cat. No.: HY-19551

Apogossypolone (ApoG2) is an orally active Bcl-2 family proteins inhibitor with K₁ values of 35, 25 and 660 nM for Bcl-2, Mcl-1 and Bcl-X₁, 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

Arctigenin

((-)-Arctigenin) Cat. No.: HY-N0035

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.

Purity: 99.69% Clinical Data: Phase 2

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg

Arglabin

((+)-Arglabin) Cat. No.: HY-16059

Arglabin ((+)-Arglabin), a natural product isolated from Artemisia glabella, is a **NLRP3 inflammasome** inhibitor. Arglabin shows anti-inflammatory and antitumor activities.



Purity: 99.49%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

ARN5187

Cat. No.: HY-103691

ARN5187 is a lysosomotropic REV-ERB β ligand with a dual inhibitory activity toward REV-ERB-mediated transcriptional regulation and autophagy. ARN5187 shows lysosomotropic potency and cytotoxicity. ARN5187 induces apoptosis.

Purity: > 98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

ARN5187 trihydrochloride

Cat. No.: HY-103691A

ARN5187 trihydrochloride is a lysosomotropic REV-ERBß ligand with a dual inhibitory activity toward REV-ERB-mediated transcriptional regulation and autophagy. ARN5187 trihydrochloride shows lysosomotropic potency and cytotoxicity. ARN5187 trihydrochloride induces apoptosis.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

AS-605240

Cat. No.: HY-10109

AS-605240 is a specific and orally active inhibitor of the PI3Ky, with an IC $_{\rm S0}$ of 8 nM, and a $\rm K_i$ of 7.8 nM.

Purity: 99.17%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg

AS1708727

AS1708727 is an orally active Foxo1 inhibitor, with EC $_{50}$ values of 0.33 μ M and 0.59 μ M for G6Pase and PEPCK, respectively.



Cat. No.: HY-123046

Purity: 99.82%

Clinical Data: No Development Reported

Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

AS1842856

Cat. No.: HY-100596

AS1842856, a specific **Foxo1** inhibitor (IC_{so} =30 nM), potently suppresses **autophagy**. AS1842856 only reduces the activity of FoxO1 by binding with it, without affecting its transcription and protein expression.

Purity: 99.83%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

Asperphenamate

Cat. No.: HY-129578

Asperphenamate, a fungal metabolite of Aspergillus flatilipes with anti-cancer effect, exhibits IC $_{50}$ values of 92.3 $\mu M,$ 96.5 μM and 97.9 μM in T47D, MDA-MB-231 and HL-60 cells, respectively.



Purity: ≥98.0%

Clinical Data: No Development Reported

Size: 5 mg, 10 mg

Aspirin

(Acetylsalicylic Acid; ASA)

Aspirin is a non-selective and irreversible inhibitor of COX-1 and COX-2 with IC $_{50}$ s of 5 and 210 $\mu g/mL$.



Cat. No.: HY-14654

Purity: 99.92% Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g

Aspirin-d3

(Acetylsalicylic Acid-d3; ASA-d3)

Aspirin-d3 (Acetylsalicylic Acid-d3) is the deuterium labeled Aspirin. Aspirin is a non-selective and irreversible inhibitor of COX-1 and COX-2 with IC $_{so}$ s of 5 and 210 $\mu g/mL$.



Cat. No.: HY-14654S

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg, 10 mg

Aspirin-d4

(Acetylsalicylic Acid-d4; ASA-d4)

Aspirin-d4 (Acetylsalicylic Acid-d4) is the deuterium labeled Aspirin. Aspirin is a non-selective and irreversible inhibitor of COX-1 and COX-2 with IC $_{so}$ s of 5 and 210 μ g/mL.

Cat. No.: HY-14654S1

Purity: 98.85%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

AT9283

AT9283 is a multi-targeted kinase inhibitor with potent activity against Aurora A/B, JAK2/3, AbI (T3151) 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.



Cat. No.: HY-50514

Purity: 99.70% Clinical Data: Phase 2

Size: 10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 50 mg, 100 mg

Ataluren

(PTC124) Cat. No.: HY-14832

Ataluren (PTC124) is an orally available CFTR-G542X nonsense allele inhibitor.

Purity: 99.71% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg

Atezolizumab

(MPDL3280A) Cat. No.: HY-P9904

Atezolizumab (MPDL3280A) is a selective humanized monoclonal IgG1 antibody against programmed death ligand 1 (PD-L1), used for cancer research.

Atezolizumab

Purity: 98.98% Clinical Data: Launched

Size: 1 mg, 5 mg, 25 mg, 50 mg

Atg4B-IN-2

Cat. No.: HY-144636

Atg4B-IN-2 is a potent competitive Atg4B inhibitor with K, value of 3.1 μM, also possesses declining PLA, inhibitory potency, IC, of 11 μM and 3.5 μM for Atg4B and PLA₂, respectively.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

ATG7-IN-1

Cat. No.: HY-145371

ATG7-IN-1 is a potent and selective inhibitor of

ATG7 ($IC_{so} = 62 \text{ nM}$).



>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Atorvastatin

Cat. No.: HY-B0589

Atorvastatin is an orally active HMG-CoA reductase inhibitor, has the ability to effectively decrease blood lipids. Atorvastatin inhibits human SV-SMC proliferation and invasion with IC_{50} s of 0.39 μM and 2.39 μM , respectively.



Purity: 99.05% Clinical Data: Launched

10 mM × 1 mL, 10 mg, 50 mg, 100 mg Size:

Atorvastatin hemicalcium salt

(CI-981; Atorvastatin hemicalcium)

Atorvastatin hemicalcium salt (CI-981) is an orally active 3-hydroxy-3-methylglutaryl coenzyme A (HMG-CoA) reductase inhibitor, has the ability to effectively decrease blood lipids.



Cat. No.: HY-17379

Purity: 99 94% Clinical Data: Launched

10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg, 500 mg

Atorvastatin-d5 hemicalcium

Cat. No.: HY-B0589S

Atorvastatin-d5 hemicalcium is the deuterium labeled Atorvastatin. Atorvastatin hemicalcium is an orally active HMG-CoA reductase inhibitor, has the ability to effectively decrease blood lipids.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Atorvastatin-d5 sodium

Cat. No.: HY-B0589S1

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 10 mg

ATRA-biotin

(Biotin-ATRA-conjugate) Cat. No.: HY-141793

ATRA-biotin (Biotin-ATRA-conjugate) is a biotin-conjugated ATRA. ATRA-biotin can be used to track ATRA in cells or a given tissue.

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Atropine sulfate monohydrate (Tropine tropate sulfate

monohydrate; DL-Hyoscyamine sulfate monohydrate) Cat. No.: HY-B0394

Atropine (Tropine tropate) sulfate monohydrate is a broad-spectrum and competitive muscarinic acetylcholine receptor (mAChR) antagonist with anti-myopia effect.

99.62% Purity: Clinical Data: Launched

10 mM × 1 mL, 100 mg Size:

Atropine-d5

(Tropine tropate-d5; DL-Hyoscyamine-d5) Cat. No.: HY-B0394S

Atropine-d5 (Tropine tropate-d5) is the deuterium labeled Atropine (sulfate monohydrate). Atropine (Tropine tropate) sulfate monohydrate is a broad-spectrum and competitive muscarinic acetylcholine receptor (mAChR) antagonist with anti-myopia effect.



Purity: >98%

Clinical Data: No Development Reported

Size 1 mg, 5 mg

Aumitin

Aumitin is a diaminopyrimidine-based autophagy inhibitor which inhibits mitochondrial respiration by targeting complex I. Aumitin inhibits starvation- and rapamycin induced autophagy dose dependently with IC $_{50}$ s of 0.12 μM and 0.24 μM ,

respectively.

Purity:

99.64%

Clinical Data: No Development Reported

5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Cat. No.: HY-124726

AUTAC1

Cat. No.: HY-134183

AUTAC1 is a MetAP2-targeting autophagy-mediated degrader (AUTAC). AUTACs contain a degradation tag and a warhead to provide target specificity. AUTAC1 contains an FBnG (p-Fluorobenzyl Guanine) and a Fumagillol moiety. Fumagillol binds covalently to MetAP2.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Autocamtide 2, amide

>98%

Clinical Data: No Development Reported

1 mg, 5 mg

AUTAC2

Purity:

Size:

Autocamtide 2, amide is a substrate (100 µM final

AUTAC2 is a FKBP12-targeting autophagy-mediated

degrader (AUTAC). AUTAC2 contains an FBnG

(p-Fluorobenzyl Guanine) and an SLF (c ligand of

FKBP) moiety. SLF binds non-covalently to FKBP12.

concentration) for CaMK family assays.

KKALRRQETVDAL-NH2

Cat. No.: HY-P1528

Cat. No.: HY-134184

Purity: 99 47%

Clinical Data: No Development Reported 1 mg, 5 mg, 10 mg

Autocamtide 2

(Autocamtide II) Cat. No.: HY-P0225

Autocamtide 2 is a highly selective peptide substrate of calcium/calmodulin-dependent protein kinase II (CaMKII). It can be used in the CaMKII activity assay.

Purity:

98 21%

Clinical Data: No Development Reported

1 mg, 5 mg

Autocamtide-2-related inhibitory peptide

Cat. No.: HY-P0214

KKALRRQETVDAL

Autocamtide-2-related inhibitory peptide is a highly specific and potent inhibitor of CaMKII with an IC₅₀ of 40 nM.

KKALRRQEAVDAL

>98% Purity:

Clinical Data: No Development Reported

Size: 1 ma, 5 ma

Autocamtide-2-related inhibitory peptide TFA

Cat. No.: HY-P0214A

Autocamtide-2-related inhibitory peptide (TFA) is a highly specific and potent inhibitor of CaMKII with an IC_{so} of 40 nM.

KKALRRQEAVDAL (TFA salt)

95.85% Purity:

Clinical Data: No Development Reported

Size: 1 ma. 5 ma

Autocamtide-2-related inhibitory peptide, myristoylated

Cat. No.: HY-P0215

Autocamtide-2-related inhibitory peptide, myristoylated is the myristoylated Autocamtide-2-related inhibitory peptide. Autocamtide-2-related inhibitory peptide is a highly specific and potent inhibitor of CaMKII

{Lys(Myr)}-KALRRQEAVDAL

with an IC₅₀ of 40 nM. >98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Autocamtide-2-related inhibitory peptide, myristoylated TFA

Cat. No.: HY-P0215A

Autocamtide-2-related inhibitory peptide, myristoylated TFA is the myristoylated Autocamtide-2-related inhibitory peptide. Autocamtide-2-related inhibitory peptide is a highly specific and potent inhibitor of CaMKII with an IC₅₀ of 40 nM.

s(Myr))-KALRRQEAVDAL (TFA salt

Purity: >98%

Clinical Data: No Development Reported

Size 1 mg, 5 mg

Autogramin-1

Cat. No.: HY-128339

Autogramin-1 potently inhibits autophagy induced by either starvation (IC $_{50}$ =0.17 μ M) or mTORC1 inhibition (Rapamycin; IC₁₀=0.44 μM).



Purity: ≥98.0%

Clinical Data: No Development Reported

Size: 5 mg, 10 mg

Autogramin-2

Cat. No.: HY-128340

Autogramin-2 potently inhibits autophagy induced by either starvation (IC $_{50}$ =0.27 μ M) or mTORC1 inhibition (Rapamycin; IC_{so}=0.14 μM).



99.82%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Autophagy inducer 2

Autophagy inducer 2 (Compound 11i) is a potent autophagy inducer. Autophagy inducer 2 exhibits apparent antiproliferative activity against the MCF-7 cell line with an IC_{50} value of 1.31 μM and remarkably inhibits the colony formation of the MCF-7 cells.

Cat. No.: HY-144637

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Autophagy inducer 4

Cat. No.: HY-146087

Autophagy inducer 4 is a Magnolol-based Mannich base derivatives, which can be used as an anticancer agent. Autophagy inducer 4 suppresses cancer cells via inducing autophagy.



Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg

Autophinib

Cat. No.: HY-101920

Autophinib is a potent, selective autophagy inhibitor with IC₅₀s of 90 nM and 40 nM for starvation- and Rapamycin-induced autophagy, respectively. Autophinib is also an ATP competitive Vacuolar Protein Sorting 34 (VPS34) inhibitor with an IC₅₀ of 19 nM.



99.56% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 50 mg, 100 mg

AZ PFKFB3 26

Cat. No.: HY-101971

AZ PFKFB3 26 is a potent and selective inhibitor of the metabolic kinase PFKFB3 with an IC_{so} of 23 nM. AZ PFKFB3 26 inhibits PFKFB1 and PFKFB2 with IC_{so} s of 2.06 and 0.384 μ M, respectively.



99.23% Purity:

Clinical Data: No Development Reported 5 mg, 10 mg, 25 mg, 50 mg Size:

AZ304

Cat. No.: HY-117273

AZ304 is an ATP-competitive dual BRAF kinase inhibitor, potently inhibits wild type BRAF, V600E mutant BRAF and wild type CRAF, with IC_{so}s of 79 nM, 38 nM and 68 nM, respectively. AZ304 also has significant effect on other kinases, such as p38 (IC₅₀, 6 nM), CSF1R (IC₅₀, 35 nM).



99.39% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 25 mg, 50 mg, 100 mg

Autophagy inducer 3

Autophagy inducer 3 has autophagy induced activity. Autophagy inducer 3 possesses robust autophagic cell death in diverse cancer cells sparing normal counterpart.



Cat. No.: HY-146052

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Autophagy-IN-C1

Cat. No.: HY-141813

Autophagy-IN-C1 not only induces apoptosis but also blocks autophagy in hepatocellular carcinoma



Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg

Avermectin B1

(Abamectin; Avermectin B1a-Avermectin B1b mixt.)

Avermectin B1 (Abamectin) is a widely used insecticide and anthelmintic. IC50 Value: N/A Target: Antiparasitic Avermectin B1 is a mixture of avermectins containing more than 80% avermectin B1a and less than 20% avermectin B1b.



Cat. No.: HY-15311

Purity: 96.89% Clinical Data: Phase 3

Size: 10 mM × 1 mL, 100 mg

AZ-PFKFB3-67

AZ-PFKFB3-67 is potent and selective metabolic kinase PFKFB3 inhibitor, with IC_{so}s of 11, 159 and 1130 nM for PFKFB3, PFKFB2 and PFKFB1

respectively.

>98% Purity:

Clinical Data: No Development Reported

Size 5 mg, 10 mg



Cat. No.: HY-101972

AZD 6482

(KIN-193)

AZD 6482 (KIN-193) is a potent and selective p110β inhibitor with an IC_{so} of 0.69 nM.



Cat. No.: HY-10344

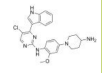
99.56% Clinical Data: Phase 1

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

AZD-3463

(ALK/IGF1R inhibitor) Cat. No.: HY-15609

AZD-3463 (ALK/IGF1R inhibitor) is an orally active ALK/IGF1R inhibitor, with a K, of 0.75 nM for ALK. AZD3463 induces apoptosis and autophagy in neuroblastoma cells.



Purity: 99 96%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

AZD-8055

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.



Cat. No.: HY-10422

99 60% Purity: Clinical Data: Phase 1

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg

AZD1208

Cat. No.: HY-15604

AZD1208 is an orally bioavailable, highly selective PIM kinases inhibitor.



Purity: 99 90% Clinical Data: Phase 1

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

AZD1208 hydrochloride

Cat. No.: HY-15604A

AZD1208 hydrochloride is an orally bioavailable, highly selective PIM kinases inhibitor.



Purity: >98% Clinical Data: Phase 1 1 mg, 5 mg

AZD7624

Cat. No.: HY-103672

AZD7624 is an inhaled p38 inhibitor, with potent anti-inflammatory activity.



98.08% Purity:

Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg

Azithromycin

(CP 62993) Cat. No.: HY-17506

Azithromycin is a macrolide antibiotic useful for the treatment of a number of bacterial infections.



≥98.0% Purity: Clinical Data: Launched

Size 10 mM × 1 mL, 50 mg, 100 mg, 200 mg, 500 mg

Azithromycin hydrate

(CP-62993 dihydrate) Cat. No.: HY-17506A

Azithromycin hydrate is a macrolide antibiotic useful for the treatment of a number of bacterial infections



>98% Purity: Clinical Data: Launched

Size 50 mg, 100 mg, 200 mg, 500 mg

Azithromycin-d3

Cat. No.: HY-17506S Azithromycin-d3 (CP 62993-d3) is the deuterium

labeled Azithromycin. Azithromycin (CP-62993) is a macrolide antibiotic useful for the treatment of a

number of bacterial infections.



>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg

Baicalin

(Baicalein 7-O-β-D-glucuronide)

Cat. No.: HY-N0197

Baicalin, as a flavonoid glycoside, is an allosteric carnitine palmityl transferase 1 (CPT1) activator. Baicalin reduces the expression of NF-κB.



99.17% Clinical Data: Launched

10 mM × 1 mL, 100 mg, 500 mg, 1 g, 5 g

Bafilomycin A1

Cat. No.: HY-100558

Bafilomycin A1 is a specific and reversible inhibitor of vacuolar H+-ATPase (V-ATPase) with IC_{so} values of 4-400 nmol/mg. Bafilomycin A1, a macrolide antibiotic, is also used as an autophagy inhibitor at the late stage.



Purity: 99.43%

Clinical Data: No Development Reported Size: 100 μg, 500 μg, 1 mg, 5 mg

Bakuchiol

((S)-(+)-Bakuchiol) Cat. No.: HY-N0235

Bakuchiol is a phytoestrogen isolated from the seeds of Psoralea corylifolia L; has anti-tumor effects.

Purity: Clinical Data: Phase 2

Size:

BAY 11-7082 is an IκBα phosphorylation and NF-κB inhibitor. BAY 11-7082 selectively and

irreversibly inhibits the TNF-α-induced phosphorylation of IkB- α , and decreases NF-kB and expression of adhesion molecules.

Purity: 99 98%

Berbamine

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

99 25%

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Bardoxolone methyl

(RTA 402; NSC 713200; CDDO Methyl ester) Cat. No.: HY-13324

Bardoxolone methyl (NSC 713200; RTA 402; CDDO Methyl ester) is a synthetic triterpenoid compound with potential antineoplastic and anti-inflammatory activities, acting as an activator of the Nrf2 pathway and an inhibitor of the NF-kB pathway.

Purity: 99 72% Clinical Data: Phase 3

10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg, 500 mg Size:

Belinostat

(PXD101; PX105684) Cat. No.: HY-10225

Belinostat (PXD101; PX105684) is a potent HDAC inhibitor with an IC₅₀ of 27 nM in HeLa cell extracts

Purity: 99 94% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg

Berbamine is a natural compound extracted from traditional Chinese medicine Barberry with anti-tumor, immunomodulatory and cardiovascular effects. Berbamine is a calcium channel

99 79% Purity: Clinical Data: Launched

Size 10 mM \times 1 mL, 5 mg, 10 mg, 20 mg

Berbamine dihydrochloride

Cat. No.: HY-N0714A

Berbamine dihydrochloride is an inhibitor of NF-κB activity with remarkable anti-myeloma efficacy.



96.62% Purity: Clinical Data: Launched

10 mM × 1 mL, 200 mg, 500 mg Size

Berberine

(Natural Yellow 18)

Berberine (Natural Yellow 18) is an alkaloid isolated from the Chinese herbal medicine Huanglian, as an antibiotic. Berberine (Natural Yellow 18) induces reactive oxygen species (ROS) generation and inhibits DNA topoisomerase.

>98% Purity: Clinical Data: Launched Size 5 mg, 10 mg, 25 mg

Berberine chloride

(Natural Yellow 18 chloride) Cat. No.: HY-18258

Berberine chloride is an alkaloid that acts as an antibiotic. Berberine chloride induces reactive oxygen species (ROS) generation and inhibits DNA topoisomerase. Antineoplastic properties.

Purity: 99.66% Clinical Data: Launched

Size 10 mM × 1 mL, 100 mg, 1 g, 5 g

Berberine chloride hydrate

(Natural Yellow 18 chloride hydrate)

Berberine chloride hydrate (Natural Yellow 18 chloride hydrate) is an alkaloid that acts as an antibiotic. Berberine chloride hydrate induces reactive oxygen species (ROS) generation and inhibits DNA topoisomerase. Antineoplastic properties.

Purity: 99.84% Clinical Data: Launched

10 mM × 1 mL, 100 mg, 1 g, 5 g



Cat. No.: HY-101273

>98.0% Purity:

Clinical Data: No Development Reported

BAR502 is a dual FXR and GPBAR1 agonist with

 IC_{so} values of 2 μ M and 0.4 μ M, respectively.

Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 50 mg, 100 mg



BAY 11-7082

BAR502

(BAY 11-7821) Cat. No.: HY-13453



Cat. No.: HY-N0714

Cat. No.: HY-N0716

Cat. No.: HY-17577

Berberine sulfate

(Natural Yellow 18 sulfate) Cat. No.: HY-N0716B

Berberine sulfate is an alkaloid isolated from the Chinese herbal medicine Huanglian, as an antibiotic. Berberine sulfate induces reactive oxygen species (ROS) generation and inhibits DNA topoisomerase. Berberine sulfate has antineoplastic properties.

Purity: >98% Clinical Data: Launched Size: 5 ma

Berberine-d6 chloride (Natural Yellow 18-d6 chloride)

Berberine-d6 (Natural Yellow 18-d6) chloride is the deuterium labeled Berberine chloride. Berberine chloride is an alkaloid that acts as an antibiotic. Berberine chloride induces reactive oxygen species (ROS) generation and inhibits DNA topoisomerase. Antineoplastic properties.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg



Cat. No.: HY-18258S

Bergapten

(5-Methoxypsoralen) Cat. No.: HY-N0370

Bergapten is a natural anti-inflammatory and anti-tumor agent. Bergapten is inhibitory towards mouse and human CYP isoforms.

Purity: 99 96% Clinical Data: Phase 3

10 mM × 1 mL, 500 mg, 1 g, 5 g Size:

Bergapten-d3

(5-Methoxypsoralen-d3)

Bergapten-d3 is deuterium labeled Bergapten. Bergapten is a natural anti-inflammatory and anti-tumor agent. Bergapten is inhibitory towards mouse and human CYP isoforms.

Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg



Cat. No.: HY-N0370S

Bergenin

(Cuscutin) Cat. No.: HY-N0017

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 Betulinic acid

(Lupatic acid; Betulic acid)

Betulinic acid is a natural pentacyclic triterpenoid, acts as a eukaryotic topoisomerase I inhibitor, with an IC_{50} of 5 μ M, and possesses anti-HIV, anti-malarial, anti-inflammatory and anti-tumor properties.

>98.0% Purity: Clinical Data: Phase 2

Size 10 mM × 1 mL, 100 mg, 200 mg, 500 mg

Cat. No.: HY-10529

Bexarotene

(LGD1069) Cat. No.: HY-14171

Bexarotene (LGD1069) is a high-affinity and selective retinoid X receptors (RXR) agonist with EC_{so}s of 33, 24, 25 nM for RXRα, RXRβ, and RXRγ, respectively. Bexarotene shows limited affinity for RAR receptors (EC $_{50}$ >10000 nM).

Purity: 99.90% Clinical Data: Launched

10 mM × 1 mL, 50 mg, 100 mg, 500 mg Size

Bexarotene D4

(LGD1069 D4) Cat. No.: HY-14171S

Bexarotene D4 is a deuterium labeled Bexarotene (LGD1069). Bexarotene (LGD1069) is a selective retinoid X receptors (RXR) agonist for the treatment of cutaneous T-cell lymphoma.

>98% Purity:

Clinical Data: No Development Reported

Size 1 mg, 5 mg

Bexarotene-d3

Cat. No.: HY-14171S1

Bexarotene-d3 (LGD1069-d3) is the deuterium labeled Bexarotene. Bexarotene (LGD1069) is a high-affinity and selective retinoid X receptors (RXR) agonist with ECsos of 33, 24, 25 nM for RXRα, RXRβ, and RXRγ, respectively.

Purity: >98% Clinical Data:

Size: 2.5 mg, 1 mg, 5 mg, 10 mg BGT226

(NVP-BGT226) Cat. No.: HY-13334A

BGT226 (NVP-BGT226) is a PI3K (with IC_{so}s of 4 nM, 63 nM and 38 nM for PI3Kα, PI3Kβ and PI3Ky)/mTOR dual inhibitor which displays potent growth-inhibitory activity against human head and neck cancer cells.

Purity: >98% Clinical Data: Phase 2 1 mg, 5 mg



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BGT226 maleate

(NVP-BGT226 maleate) Cat. No.: HY-13334

BGT226 (NVP-BGT226 maleate) is a PI3K (with IC $_{s0}$ 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.



Purity: 99.73% Clinical Data: Phase 2

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

BIA 10-2474-d3

Purity:

Size:

BI-D1870

BIA 10-2474-d3 is the deuterium labeled BIA 10-2474. BIA 10-2474 is an inhibitor of fatty acid amide hydrolase (FAAH) with $\rm IC_{50}$ values of 50 to 70mg/kg in various rat brain regions.

BI-D1870 is an ATP-competitive, cell permeable and

10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 50 mg

brain penetrated inhibitor of RSK isoforms, with

IC₅₀s of 31 nM/24 nM/18 nM/15 nM for RSK1/RSK2/RSK3/RSK4, respectively.

99 14%

Clinical Data: No Development Reported



Cat. No.: HY-14249S

Cat. No.: HY-19740S

Cat. No.: HY-10510

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

BIA 10-2474

Cat. No.: HY-19740

BIA 10-2474 is an inhibitor of fatty acid amide hydrolase (FAAH) with $\rm IC_{50}$ values of 50 to 70mg/kg in various rat brain regions.

Purity: 98.41%

Clinical Data: No Development Reported
Size: 5 mg, 10 mg, 50 mg, 100 mg

Bicalutamide

Cat. No.: HY-14249

Bicalutamide is an orally active non-steroidal androgen receptor (AR) antagonist. Bicalutamide can be used for the research of prostate cancer.

Purity: 99.62% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 200 mg, 500 mg, 1 g, 5 g

Bicalutamide-d4

Bicalutamide-d4 is the deuterium labeled Bicalutamide. Bicalutamide is an orally active non-steroidal androgen receptor (AR) antagonist. Bicalutamide can be used for the research of prostate cancer.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 10 mg

Bicyclol

(SY801) Cat. No.: HY-B0766

Bicyclol(SY 801) is a anti-hepatitis drug. Target: HBV Oral administration of bicyclol normalized the elevated serum transaminases (ALT, AST) by approximately 50% in chronic viral hepatitis B and C, and also showed certain level of inhibiting HBV and HCV replication.



Purity: 99.84% Clinical Data: Launched

Size: $10 \text{ mM} \times 1 \text{ mL}$, 5 mg, 10 mg, 25 mg, 50 mg

Bigelovin

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.

Purity: 99.81%

Clinical Data: No Development Reported

Size: 5 mg, 10 mg

H H O

Cat. No.: HY-116506

BIIB021

(CNF2024) Cat. No.: HY-10212

BIIB021 (CNF2024) is an orally active, fully synthetic inhibitor of HSP90 with a K_i and an EC_{sn} of 1.7 nM and 38 nM, respectively.

Purity: 99.93% Clinical Data: Phase 2

Size: $10 \text{ mM} \times 1 \text{ mL}$, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

Bilobalide

((-)-Bilobalide)

Bilobalide, a sesquiterpene trilactone constituent of Ginkgo biloba, inhibits the NMDA-induced efflux of choline with an IC $_{50}$ 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.

Purity: ≥98.0%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg



Cat. No.: HY-N0076

Binimetinib

(MEK162; ARRY-162; ARRY-438162)

Binimetinib (MEK162) is an oral and selective MEK1/2 inhibitor. Binimetinib (MEK162) inhibits MEK with an $\rm IC_{50}$ of 12 nM.

N H O OH

Cat. No.: HY-15202

Purity: 99.24% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg

Binimetinib-13C,d3

(MEK162-13C,d3; ARRY-162-13C,d3; ARRY-438162-13C,d3) Cat. No.: HY-15202S

Binimetinib-13C,d3 (MEK162-13C,d3) is the 13C- and deuterium labeled Binimetinib. Binimetinib (MEK162) is an oral and selective MEK1/2 inhibitor. Binimetinib (MEK162) inhibits MEK with an IC_{50} of 12 nM.

NN NH NH NO OH

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Biochanin A

(4-Methylgenistein; Olmelin)

Biochanin A is a naturally occurring fatty acid amide hydrolase (FAAH) inhibitor, which inhibits FAAH with IC $_{50}$ S of 1.8, 1.4 and 2.4 μ M for mouse, rat, and human FAAH, respectively.

Cat. No.: HY-14595

Purity: 98.98%

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 200 mg, 500 mg

Bisdemethoxycurcumin

(Curcumin III; Didemethoxycurcumin)

Bisdemethoxycurcumin(Curcumin III; Didemethoxycurcumin) is a natural derivative of curcumin with anti-inflammatory and anti-cancer activities.

HO CHICA

Cat. No.: HY-N0007

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)

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.

Cat. No.: HY-N0007S

Purity: >98%

Clinical Data: No Development Reported

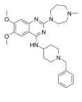
Size: 1 mg, 5 mg

BIX-01294

BIX-01294 is a reversible and highly selective G9a and GLP Histone Methyltransferase inhibitor, with IC $_{s0}s$ of of 1.7 μM and 0.9 $\mu\text{M},$ respectively.

Purity: 99.59%

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 10 mg, 50 mg



Cat. No.: HY-10587

BIX-01294 trihydrochloride

BIX-01294 trihydrochloride is a reversible and highly selective **G9a and GLP Histone**

Methyltransferase inhibitor, with IC_{s0} s of of 1.7 μ M and 0.9 μ M, respectively.

HN (N)

Cat. No.: HY-108239

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

BL-918

BL-918 is an orally active UNC-51-like kinase 1 (ULKI) activator with an EC $_{50}$ of 24.14 nM. BL-918 exerts its cytoprotective **autophagic** effect by targeting ULK complex. BL-918 has the potential for Parkinson's disease (PD) treatment.



Cat. No.: HY-124729

Purity: 98.36%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

BMS-582949 hydrochloride

Cat. No.: HY-14305A

BMS-582949 hydrochloride is an orally active and highly selective $p38\alpha$ MAPK inhibitor, with an IC_{s0} of 13 nM. BMS-582949 hydrochloride displays a significantly improved pharmacokinetic profile and is effective in inflammatory disease.



Purity: 98.29% Clinical Data: Phase 2

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Bortezomib

(PS-341; LDP-341; NSC 681239)

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. Bortezomib disrupts the cell cycle, induces apoptosis, and inhibits NF- κ B.



Cat. No.: HY-10227

Purity: 99.97% Clinical Data: Launched

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

Bortezomib-d8

(PS-341-d8; LDP-341-d8; NSC 681239-d8)

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.

Cat. No.: HY-10227S

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Bosutinib D8

(SKI-606 D8) Cat. No.: HY-10158S

Bosutinib D8 (SKI-606 D8) is a deuterium labeled Bosutinib. Bosutinib is a dual Src/Abl inhibitor with IC₅₀s of 1.2 nM and 1 nM, respectively.



Purity: >99.0%

Clinical Data: No Development Reported

Size

Brazilin

Cat. No.: HY-N0072

Brazilin is a red dye precursor obtained from the heartwood of several species of tropical hardwoods. Brazilin inhibits the cells proliferation, promotes apoptosis, and induces autophagy through the AMPK/mTOR pathway.

99.26% Purity:

Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 20 mg

BRD5631

Cat. No.: HY-125197

BRD5631 is an autophagy enhancer, enhances autophagy through an mTOR-independent pathway.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Brevilin A

Cat. No.: HY-N2959

Brevilin A is a sesquiterpene lactone isolated from Centipeda minima with anti-tumor activity. Brevilin A is a selective inhibitor of JAK-STAT signal pathway by attenuating the JAKs activity and blocking STAT3 signaling (IC $_{50}$ = 10.6 μ M) in Cancer Cells.

Purity: 99.77%

Clinical Data: No Development Reported

Size: 5 mg, 10 mg

Bosutinib

(SKI-606) Cat. No.: HY-10158

Bosutinib is a dual Src/Abl inhibitor with IC_{so}s of 1.2 nM and 1 nM, respectively.



99 96% Purity: Clinical Data: Launched

10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg

BPO-27 racemate

Cat. No.: HY-19778A

BPO-27 racemate is a potent CFTR inhibitor with an IC_{50} of 8 nM.



Purity: 98.37%

Clinical Data: No Development Reported 1 mg, 5 mg, 10 mg

BRD4/CK2-IN-1

Cat. No.: HY-145260

BRD4/CK2-IN-1 is the first highly effective and oral active dual-target inhibitor of BRD4/CK2 (bromodomain-containing protein 4/casein kinase 2), with IC_{so}s of 180 nM and 230 nM for BRD4 and CK2, respectively.



>98% Purity:

Clinical Data: No Development Reported

Size 1 mg, 5 mg

Brefeldin A

(BFA; Cyanein; Decumbin)

Brefeldin A (BFA) is a lactone antibiotic and a specific inhibitor of protein trafficking. Brefeldin A blocks the transport of secreted and membrane proteins from endoplasmic reticulum to Golgi apparatus. Brefeldin A is also an autophagy and



Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg Size:

Cat. No.: HY-16592

Briciclib

(ON 014185)

Briciclib (ON 014185) is a derivative of ON 013100, and has the potential in targeting eIF4E for solid cancers.



Cat. No.: HY-16366

99.65% Purity: Clinical Data: Phase 1

10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 50 mg, 100 mg

Britannin

Cat. No.: HY-N3005

Britannin, isolated from Inula aucheriana, is a sesquiterpene lactone. Britannin induces apoptosis and autophagy by activating AMPK regulated by ROS in liver cancer cells. Britannin has anti-proliferative and anti-inflammatory activities



Purity: 99 90%

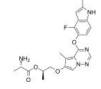
Clinical Data: No Development Reported

Size: 5 mg, 10 mg

Brivanib (alaninate)

(BMS-582664) Cat. No.: HY-10336

Brivanib alaninate (BMS-582664) is an ATP-competitive inhibitor against VEGFR2 with an IC_{so} of 25 nM; has moderate potency against VEGFR-1 and FGFR-1, but more than 240-fold against PDGFRß.



Purity: 99.45% Clinical Data: Phase 3

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Bromhexine hydrochloride

99 24%

Clinical Data: Phase 3

Brivanib

Purity:

(BMS-540215)

Cat. No.: HY-B0372A

Cat. No.: HY-10337

Bromhexine hydrochloride is a potent and specific TMPRSS2 protease inhibitor with an IC₅₀ of 0.75 μM. Bromhexine hydrochloride can prevent and manage SARS-CoV-2 infection. Bromhexine hydrochloride is an autophagy agonist.

Brivanib (BMS-540215) is an ATP-competitive

inhibitor against VEGFR2 with an IC₅₀ of 25 nM,

and has moderate potency against VEGFR-1 and FGFR-1, but >240-fold against PDGFR-β.

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg



Purity: 99.39% Clinical Data: Launched

10 mM × 1 mL, 500 mg, 5 g, 10 g

Bromhexine-d3 hydrochloride

Cat. No.: HY-B0372AS

Bromhexine-d3 (hydrochloride) is deuterium labeled Bromhexine (hydrochloride). Bromhexine hydrochloride is a potent and specific TMPRSS2 protease inhibitor with an IC50 of 0.75 μ M. Bromhexine hydrochloride can prevent and manage SARS-CoV-2 infection.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 ma, 5 ma

Bromocriptine mesylate

(CB-154) Cat. No.: HY-12705A

Bromocriptine mesylate is a potent dopamine D2/D3 receptor agonist, which binds D2 dopamine receptor with pK, of 8.05±0.2.



99.98% Purity: Clinical Data: Launched

 $10~\text{mM}\times1~\text{mL},\,50~\text{mg},\,100~\text{mg},\,200~\text{mg}$ Size

Butein

(2',3,4,4'-tetrahydroxy Chalcone) Cat. No.: HY-16558

Butein is a cAMP-specific PDE inhibitor with an IC_{so} of 10.4 μM for PDE4. Butein is a specific protein tyrosine kinase inhibitor with IC₅₀s of 16 and 65 µM for EGFR and p60^{c-src} in HepG2 cells.

99.95% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 50 mg, 100 mg Size

BX795

Cat. No.: HY-10514

BX795 is a potent and selective inhibitor of PDK1, with an IC₅₀ of 6 nM. BX795 is also a potent and relatively specific inhibitor of TBK1 and IKKE, with an IC_{so} of 6 and 41 nM, respectively.



99.17% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg, 200 mg

C2 Ceramide

(Ceramide 2) Cat. No.: HY-101180

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).



Purity: ≥98.0%

Clinical Data: No Development Reported

5 mg Size:

C646

Cat. No.: HY-13823

C646 is a selective and competitive histone acetyltransferase p300 inhibitor with K, of 400 nM, and is less potent for other acetyltransferases.



≥98.0%

Clinical Data: No Development Reported 10 mM × 1 mL, 10 mg, 50 mg

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C8-Ceramide

(N-Octanoyl-D-erythro-sphingosine)

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.



Cat. No.: HY-108391

Purity: >98.0%

Clinical Data: No Development Reported

Size: 5 mg

Cabazitaxel

Purity:

Size:

CA-5f

CA-5f is a potent late-stage

production. Anti-tumor activity.

99 40%

Clinical Data: No Development Reported

(XRP6258; RPR-116258A; taxoid XRP6258)

macroautophagy/autophagy inhibitor via

and SQSTM1 protein, and also increases ROS

inhibiting autophagosome-lysosome fusion. CA-5f increases LC3B-II (a marker to monitor autophagy)

Cabazitaxel is a semi-synthetic derivative of the natural taxoid 10-deacetylbaccatin III with potential antineoplastic activity.



Cat. No.: HY-15459S1

Cat. No.: HY-15459

Cat. No.: HY-112698

Purity: 99 96% Clinical Data: Launched

Cabazitaxel-d9

10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 500 mg

(XRP6258-d9; RPR-116258A-d9; taxoid XRP6258-d9)

Cabazitaxel-d9 is deuterium labeled Cabazitaxel.

Cabazitaxel is a semi-synthetic derivative of the natural taxoid 10-deacetylbaccatin III with

potential antineoplastic activity.

>98%

Clinical Data: No Development Reported

1 mg, 5 mg

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

CA77.1

Cat. No.: HY-134923

CA77.1 is a potent, brain-penetrant and orally active chaperone-mediated autophagy (CMA) activator with favorable pharmacokinetics. CA77.1 is a derivative of AR7 (HY-101106) and can increase the expression of the lysosomal receptor LAMP2A in lysosomes.

Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg Size:

Cabazitaxel-d6

(XRP6258-d6; RPR-116258A-d6; taxoid XRP6258-d6) Cat. No.: HY-15459S

Cabazitaxel-d6 (XRP6258-d6) is the deuterium labeled Cabazitaxel. Cabazitaxel is a semi-synthetic derivative of the natural taxoid 10-deacetylbaccatin III with potential antineoplastic activity.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 10 mg

Cabergoline

(FCE-21336) Cat. No.: HY-15296

Cabergoline is an ergot derived-dopamine D₂-like receptor agonist that has high affinity for D₂, D_{2} , and 5-HT₂₈ receptors (K_{i} =0.7, 1.5, and 1.2, respectively).



99.80% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg

Cabergoline-d5

Purity:

Size:

(FCE-21336-d5) Cat. No.: HY-15296S

Cabergoline-d5 (FCE-21336-d5) is the deuterium labeled Cabergoline. Cabergoline is an ergot derived-dopamine D₂-like receptor agonist that has high affinity for D_{2} , D_{3} , and $5-HT_{2B}$ receptors (K_i=0.7, 1.5, and 1.2, respectively).



Purity: >98%

Clinical Data: No Development Reported

Size 2.5 mg

Calcimycin

(A-23187; Antibiotic A-23187)

Calcimycin (A-23187) is an antibiotic and a unique divalent cation ionophore (like calcium and magnesium). Calcimycin induces Ca²⁺-dependent cell death by increasing intracellular calcium concentration. Calcimycin inhibits the growth of Gram-positive bacteria and some fungi.



Cat. No.: HY-N6687

Purity: 99.56% Clinical Data: Phase 3

 $10 \text{ mM} \times 1 \text{ mL}, 1 \text{ mg}, 5 \text{ mg}$

Cabergoline-d6

(FCE-21336-d6) Cat. No.: HY-15296S1

Cabergoline-d6 is deuterium labeled Cabergoline. Cabergoline is an ergot derived-dopamine D2-like receptor agonist that has high affinity for D2, D3, and 5-HT2B receptors (Ki=0.7, 1.5, and 1.2, respectively).



Purity: >98%

Clinical Data:

Size 1 mg, 5 mg

Calcimycin hemicalcium salt (A-23187 hemicalcium salt;

Antibiotic A-23187 hemicalcium salt) Cat. No.: HY-N6687A

Calcimycin hemicalcium salt (A-23187 hemicalcium salt) is an antibiotic and a unique divalent cation ionophore (like calcium and magnesium). Calcimycin hemicalcium salt induces Ca²⁺-dependent cell death by increasing intracellular calcium concentration.



Purity: >98%

Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg

Calcimycin hemimagnesium

(A-23187 hemimagnesium; Antibiotic A-23187 hemimagnesium: No.: HY-N6687B

Calcimycin (A-23187) hemimagnesium is an antibiotic and a unique divalent cation ionophore (like calcium and magnesium). Calcimycin hemimagnesium induces Ca2+-dependent cell death by increasing intracellular calcium concentration.



>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Calcineurin substrate

Cat. No.: HY-P0228

Calcineurin substrate is a peptide from the regulatory RII subunit of cAMP-dependent protein kinase. It can be used in the calcineurin activity assay.

DLDVPIPGREDRRVSVAAE

Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg

Calcineurin substrate TFA

Cat. No.: HY-P0228A

Calcineurin substrate (TFA) is a peptide from the regulatory RII subunit of cAMP-dependent protein kinase. Calcineurin substrate (TFA) can be used in the calcineurin activity assay.

DLDVPIPGRFDRRVSVAAE (TFA salt)

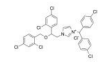
Purity: 99 57%

Clinical Data: No Development Reported 500 μg, 1 mg, 5 mg

Calmidazolium chloride

(R 24571) Cat. No.: HY-103319

Calmidazolium chloride (R 24571) is a calmodulin (CaMK) antagonist, antagonizing CaM-dependent phosphodiesterase and calmodulin-induced activation of erythrocyte Ca2+-transporting ATPase with IC₅₀s of 0.15 and 0.35 μM, respectively.



Purity: 98.93%

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

Calmodulin-Dependent Protein Kinase II (290-309)

Cat. No.: HY-P1479

Calmodulin-Dependent Protein Kinase II (290-309) is a potent CaMK antagonist with an IC₅₀ of 52 nM for inhibition of Ca2+/calmodulin-dependent protein kinase II.

LKKFNARRKLKGAILTTMLA

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

CaMKII-IN-1

Calmodulin-Dependent Protein Kinase II(290-309) acetate

Cat. No.: HY-P1479A

Calmodulin-Dependent Protein Kinase II (290-309) acetate is a potent CaMK antagonist with an IC_{50} of 52 nM for inhibition of Ca2+/calmodulin-dependent protein kinase II.

CaMKII-IN-1 is a potent and highly selective CaMKII inhibitor with IC50 of 63 nM; significantly high selectivity against CaMKIV, MLCK, p38a, Akt1, and PKC. IC50 value: 63 nM Target: CaMKII.



Cat. No.: HY-18271

99.74% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

98.97%

Capivasertib

Purity:

(AZD5363) Cat. No.: HY-15431

Capivasertib (AZD5363) is an orally active and potent pan-AKT kinase inhibitor with ICso of 3, 7 and 7 nM for Akt1,Akt2 and Akt3, respectively.



Purity: 99.83% Clinical Data: Phase 3

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Capsaicin

((E)-Capsaicin) Cat. No.: HY-10448

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.



99.85% Purity: Clinical Data: Launched

10 mM × 1 mL, 50 mg, 100 mg

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Capsaicin-d3

((E)-Capsaicin-d3) Cat. No.: HY-10448S1

Capsaicin-d3 ((E)-Capsaicin-d3) is the deuterium labeled Capsaicin. Capsaicin ((E)-Capsaicin), an active component of chili peppers, is a $\mathsf{TRPV1}$ agonist. Capsaicin has pain relief, antioxidant, anti-inflammatory, neuroprotection and anti-cancer effects.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 10 mg

Carbamazepine

(CBZ; NSC 169864) Cat. No.: HY-B0246

Carbamazepine, a sodium channel blocker, is an anticonvulsant drug.

Purity: 99 90% Clinical Data: Launched

10 mM × 1 mL, 100 mg, 500 mg

Capsaicinoid

Capsaicinoid is a mixture of Capsaicin and Dihydrocapsaicin. Capsaicinoid is an capsaicin receptor (TRPV1) agonist.



Cat. No.: HY-10448A

Purity: 99 46%

Clinical Data: No Development Reported Size: 10 mM × 1 mL, 50 mg

Carbamazepine-d10

(CBZ-d10; NSC 169864-d10)

Carbamazepine-D10 (CBZ-d10) is the deuterium labeled Carbamazepine. Carbamazepine (CBZ), a sodium channel blocker, is an anticonvulsant agent.



Cat. No.: HY-B0246S

Purity: >98%

Clinical Data: No Development Reported

2.5 mg, 1 mg

Carbamazepine-d2

(CBZ-d2; NSC 169864-d2) Cat. No.: HY-B0246S1

Carbamazepine-d2 (CBZ-d2) is the deuterium labeled Carbamazepine. Carbamazepine, a sodium channel blocker, is an anticonvulsant drug.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 10 mg

Carboplatin (NSC 241240)

Cat. No.: HY-17393

Carboplatin (NSC 241240) is a DNA synthesis inhibitor which binds to DNA, inhibits replication and transcription and induces cell death. Carboplatin (NSC 241240) is a derivative of CDDP and a potent anti-cancer agent.



99.96% Purity: Clinical Data: Launched

Size 100 mg, 200 mg, 500 mg

Carboplatin-d4

(NSC 241240-d4) Cat. No.: HY-17393S

Carboplatin-d4 (NSC 241240-d4) is the deuterium labeled Carboplatin. Carboplatin (NSC 241240) is a DNA synthesis inhibitor which binds to DNA, inhibits replication and transcription and induces cell death. Carboplatin (NSC 241240) is a derivative of CDDP and a potent anti-cancer agent.



Purity:

Clinical Data: No Development Reported

Size: 1 ma, 5 ma

Carfilzomib

(PR-171) Cat. No.: HY-10455

Carfilzomib (PR-171) is an irreversible proteasome inhibitor with an IC₅₀ of 5 nM in ANBL-6 and RPMI 8226 cells.



99.96% Purity: Clinical Data: Launched

Size 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

Carfilzomib-d8

Cat. No.: HY-10455S

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: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Carprofen

Carprofen is a nonsteroid anti-inflammatory agent,

acts as a multi-target FAAH/COX inhibitor, with IC_{so}s of 3.9 μM, 22.3 μM and 78.6 μM for COX-2, COX-1 and FAAH, respectively.



Cat. No.: HY-B1227

Purity: 99.96% Clinical Data: Launched

10 mM × 1 mL, 100 mg

Carprofen-d3

Cat. No.: HY-B1227S

Carprofen-d3 is the deuterium labeled Carprofen. Carprofen is a nonsteroid anti-inflammatory agent, acts as a multi-target FAAH/COX inhibitor, with IC_{so}s of 3.9 μM, 22.3 μM and 78.6 μM for COX-2, COX-1 and FAAH, respectively.

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 10 mg

Carvedilol-d3

Clinical Data: Launched

Carvedilol

(BM 14190)

heart failure.

Purity:

Size:

AA is the deuterium labeled Carvedilol. Carvedilol (BM 14190) is a non-selective $\beta/\alpha-1$ blocker.

10 mM × 1 mL, 100 mg, 500 mg

Carvedilol inhibits lipid peroxidation in a dose-dependent manner with an IC $_{50}$ of 5 $\mu M.$

Carvedilol (BM 14190) is a non-selective $\beta/\alpha-1$

blocker. Carvedilol inhibits lipid peroxidation in

a dose-dependent manner with an IC_{50} of 5 μ M.

Carvedilol is a multiple action antihypertensive

99 87%

agent with potential use in angina and congestive

Purity: >98%

Clinical Data: No Development Reported

1 mg, 10 mg

Carvedilol phosphate hemihydrate

(BM 14190 phosphate hemihydrate)

Carvedilol phosphate hemihydrate (BM 14190 phosphate hemihydrate) is a non-selective $\beta/\alpha-1$ blocker. Carvedilol phosphate hemihydrate inhibits lipid peroxidation with an IC_{50} of 5 μ M.

Cat. No.: HY-B0006A

Purity: >98% Clinical Data: Launched 1 mg, 5 mg

Carvedilol-d4

(BM 14190-d4) Cat. No.: HY-B0006S1

Carvedilol-d4 (BM 14190-d4) is the deuterium labeled Carvedilol. Carvedilol (BM 14190) is a non-selective $\beta/\alpha-1$ blocker. Carvedilol inhibits lipid peroxidation in a dose-dependent manner with an IC $_{50}$ of 5 $\mu M.$



>98% Purity:

Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg

Carvedilol-d5

(BM 14190-d5) Cat. No.: HY-B0006S2

Carvedilol-d5 is deuterium labeled Carvedilol. Carvedilol (BM 14190) is a non-selective $\beta/\alpha-1$ blocker. Carvedilol inhibits lipid peroxidation in a dose-dependent manner with an IC50 of 5 $\mu M_{\rm \cdot}$



Cat. No.: HY-B0006

Cat. No.: HY-B0006S

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

CCT007093

Cat. No.: HY-15880

CCT007093 is an effective protein phosphatase 1D (PPM1D Wip1) inhibitor. Wip1 inhibition can activate the mTORC1 pathway and enhance hepatocyte proliferation after hepatectomy.

Purity: >98.0%

Clinical Data: No Development Reported

10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 500 mg Size

CCT020312

Cat. No.: HY-119240

CCT020312 is a selective EIF2AK3/PERK activator. CCT020312 elicits EIF2A phosphorylation in cells.



98.97% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

CCT128930

Cat. No.: HY-13260

CCT128930 is a ATP-competitive and selective inhibitor of AKT (IC₅₀=6 nM for AKT2).



Purity: 99.69%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

CCT128930 hydrochloride

Cat. No.: HY-13260A

CCT128930 hydrochloride is a potent and selective inhibitor of AKT (IC₅₀=6 nM).



98.32%

Clinical Data: No Development Reported 5 mg, 10 mg, 50 mg, 100 mg Size:

CD437

Cearoin

(AHPN) Cat. No.: HY-100532

CD437 is a selective Retinoic Acid Receptor y (RARy) agonist.



>98.0% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Cat. No.: HY-N8418

Cearoin increases autophagy and apoptosis through the production of ROS and the activation of ERK.

Purity: > 98.0%

Clinical Data: No Development Reported

Cediranib maleate

(AZD-2171 maleate) Cat. No.: HY-13049

Cediranib maleate (AZD-2171 maleate) is a highly potent, orally available VEGFR inhibitor with IC_{so}s of <1, <3, 5, 5, 36, 2 nM for Flt1, KDR, Flt4, PDGFRα, PDGFRβ, c-Kit, respectively.



99.74% Purity: Clinical Data: Phase 3

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

CFTR(inh)-172

Cat. No.: HY-16671

CFTR(inh)-172 is a potent and selective blocker of the CFTR chloride channel; reversibly inhibits CFTR short-circuit current in less than 2 minutes with a K, of 300 nM.



98.70% Purity:

Clinical Data: No Development Reported 10 mM \times 1 mL, 5 mg, 10 mg, 50 mg Size

cGMP-HTL

(cGMP-HaloTag-ligand) Cat. No.: HY-133869

cGMP-HTL contains a HT-ligand, a linker and the Cys-S-cGMP (autophagy tag). cGMP-HTL increases the K63-linked ubiquitination of mitochondria.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

CD437-13C6

(AHPN-13C6) Cat. No.: HY-100532S

CD437-13C6 is the 13C- and deuterium labeled. CD437 is a selective Retinoic Acid Receptor v (RARy) agonist.



>98% Purity:

Clinical Data:

Size: 1 mg, 5 mg

Cediranib

(AZD2171) Cat. No.: HY-10205

Cediranib (AZD2171) is a highly potent, orally available VEGFR tyrosine kinase inhibitor with IC_{so}s of <1, <3, 5, 5, 36, 2 nM for Flt1, KDR, Flt4, PDGFRα, PDGFRβ, c-Kit, respectively.



Purity: 99.58% Clinical Data: Phase 3

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

CFTR corrector 2

Cat. No.: HY-125381

CFTR corrector 2 is a cystic fibrosis transmembrane conductance corrector (CFTR), extracted from patent US20140274933.



98.29% Purity: Clinical Data: Phase 2

Size 5 mg, 10 mg, 50 mg, 100 mg

CGI-1746

Cat. No.: HY-11999

CGI-1746 is a potent and highly selective

inhibitor of

the Btk with IC_{50} of 1.9

Purity: 98.01%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Ch55-O-C3-NH2

(RAR ligand 1)

Cat. No.: HY-111843

Ch55-O-C3-NH2 (RAR ligand 1) is a Ch 55-based ligand, which targets RAR. Ch55-O-C3-NH2 (RAR ligand 1) binds to cIAP1 ligand Bestatin via a linker to form SNIPER.



Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg

Chelerythrine

Cat. No.: HY-N2359

Chelerythrine is a natural alkaloid, acts as a potent and selective Ca²+/phospholopid-dependent PKC antagonist, with an IC $_{50}$ of 0.7 μ M. Chelerythrine has antitumor, antidiabetic and anti-inflammatory activity.



Purity: > 98%

Clinical Data: No Development Reported Size: 5 mg, 10 mg, 20 mg

Chelerythrine chloride

Chelerythrine chloride is a potent, cell-permeable inhibitor of **protein kinase** C, with an IC $_{50}$ of 660 nM. Chelerythrine chloride inhibits the **Bcl-XL-Bak BH3** peptide binding with IC $_{50}$ of 1.5 μ M and displaces Bax from Bcl-XL. Chelerythrine chloride induces **apoptosis** and **autophagy**.



Cat. No.: HY-12048

Purity: 98.56%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Chenodeoxycholic Acid

(CDCA) Cat. No.: HY-76847

Chenodeoxycholic Acid is a hydrophobic primary bile acid that activates nuclear receptors (FXR) involved in cholesterol metabolism.

Purity: ≥98.0%
Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg

Chenodeoxycholic acid-13C

(CDCA-13C) Cat. No.: HY-76847S2

Chenodeoxycholic acid-13C (CDCA-13C) is the 13C-labeled Chenodeoxycholic Acid.
Chenodeoxycholic Acid is a hydrophobic primary bile acid that activates nuclear receptors (FXR) involved in cholesterol metabolism.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Chenodeoxycholic Acid-d4

(CDCA-d4) Cat. No.: HY-76847S

Chenodeoxycholic Acid-d4 (CDCA-d4) is the deuterium labeled Chenodeoxycholic Acid. Chenodeoxycholic Acid is a hydrophobic primary bile acid that activates nuclear receptors (FXR) involved in cholesterol metabolism.

Purity: >98%

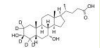
Clinical Data: No Development Reported

Size: 5 mg, 50 mg

Chenodeoxycholic acid-d5

(CDCA-d5) Cat. No.: HY-76847S3

Chenodeoxycholic acid-d5 (CDCA-d5) is the deuterium labeled Chenodeoxycholic Acid. Chenodeoxycholic Acid is a hydrophobic primary bile acid that activates nuclear receptors (FXR) involved in cholesterol metabolism.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Chloroquine

Cat. No.: HY-17589A

Chloroquine is an antimalarial and anti-inflammatory agent widely used to treat malaria and rheumatoid arthritis. Chloroquine is an autophagy and toll-like receptors (TLRs) inhibitor.

Purity: 99.50% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 200 mg, 500 mg

Chloroquine dihydrochloride

Cat. No.: HY-17589B

Chloroquine dihydrochloride is an antimalarial and anti-inflammatory agent widely used to treat malaria and rheumatoid arthritis. Chloroquine dihydrochloride is an autophagy and toll-like receptors (TLRs) inhibitor.

Purity: >98%
Clinical Data: Launched
Size: 1 mg, 5 mg

Chloroquine phosphate

Cat. No.: HY-17589

Chloroquine phosphate is an antimalarial and anti-inflammatory agent widely used to treat malaria and rheumatoid arthritis. Chloroquine phosphate is an autophagy and toll-like receptors (TLRs) inhibitor.

Purity: 99.89% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 200 mg, 500 mg

Chloroquine-d4 phosphate

Cat. No.: HY-17589S1

Chloroquine-d4 phosphate is the deuterium labeled Chloroquine phosphate. Chloroquine phosphate is an antimalarial and anti-inflammatory agent widely used to treat malaria and rheumatoid arthritis. Chloroquine phosphate is an autophagy and toll-like receptors (TLRs) inhibitor.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

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

Chloroquine-d5

Cat. No.: HY-17589AS

Chloroquine D5 is deuterium labeled Chloroquine. Chloroquine is an antimalarial and anti-inflammatory agent widely used to treat malaria and rheumatoid arthritis. Chloroquine is an autophagy and toll-like receptors (TLRs) inhibitor.

Purity: >98.0%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Ciclopirox

(HOE296b) Cat. No.: HY-B0450

Ciclopirox (HOE296b) is a synthetic antifungal agent that can be used for superficial mycoses reseaech.

Purity: 99 75% Clinical Data: Launched

10 mM × 1 mL, 50 mg, 100 mg

Ciclopirox-d11

Purity:

Size:

(HOE296b-d11)

Ciclopirox-d11 (HOE296b-d11) is the deuterium labeled Ciclopirox. Ciclopirox (HOE296b) is a synthetic antifungal agent that can be used for superficial mycoses reseaech.

Chloroquine-d5 diphosphate

phosphate is an antimalarial and

>98%

Clinical Data: No Development Reported

1 mg, 5 mg

Chloroquine-d5 diphosphate is the deuterium

labeled Chloroquine (phosphate). Chloroquine

anti-inflammatory agent widely used to treat malaria and rheumatoid arthritis.



Cat. No.: HY-B0450S

Cat. No.: HY-17589S

Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg

Ciclopirox-d11 sodium

Cat. No.: HY-B0450S1

Ciclopirox-d11 (sodium) is deuterium labeled Ciclopirox. Ciclopirox (HOE296b) is a synthetic antifungal agent that can be used for superficial mycoses reseaech.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg Cilengitide (EMD 121974)

Cilengitide (EMD 121974) is a potent and selective inhibitor of the integrins $\alpha_{\nu}\beta_{3}$ and $\alpha_{\nu}\beta_{5}$. Cilengitide inhibits binding of isolated α, β . and $\alpha_{\nu}\beta_{\nu}$ to Vitronectin with an IC₅₀ value of 4 and 79 nM, respectively.



Cat. No.: HY-16141

99 32% Purity: Clinical Data: Phase 3

Size 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Cilengitide TFA

(EMD 121974 TFA)

Cilengitide is a potent and selective integrin inhibitor for $\alpha_s \beta_s$ and $\alpha_s \beta_s$ receptor, with IC₅₀ values of 4 nM and 79 nM, respectively.



Cat. No.: HY-16143

98.85% Purity:

Clinical Data: No Development Reported 10 mM \times 1 mL, 5 mg, 10 mg, 25 mg Size

Cilofexor

(GS-9674) Cat. No.: HY-109083

Cilofexor (GS-9674) is a potent, selective and orally active nonsteroidal FXR agonist with an EC₅₀ of 43 nM. Cilofexor has anti-inflammatory and antifibrotic effects. Cilofexor has the potential for primary sclerosing cholangitis (PSC) and nonalcoholic steatohepatitis (NASH) research.



Purity: 99.82% Clinical Data: Phase 3

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg Size

Cilostazol

(OPC 13013) Cat. No.: HY-17464

Cilostazol (OPC 13013) is a potent and selective inhibitor of phosphodiesterase (PDE) 3A, the isoform of PDE 3 in the cardiovascular system, with an IC_{50} of 0.2 μ M.



Purity: 99.91% Clinical Data: Launched

Size 10 mM × 1 mL, 50 mg, 100 mg

Cilostazol-d11

Cat. No.: HY-17464S

Cilostazol-d11 is the deuterium labeled Cilostazol. Cilostazol (OPC 13013) is a potent and selective inhibitor of phosphodiesterase (PDE) 3A, the isoform of PDE 3 in the cardiovascular system, with an IC_{50} of 0.2 μ M.



Purity: >98%

Clinical Data: No Development Reported

1 mg, 10 mg

Cilostazol-d4

(OPC-13013-d4) Cat. No.: HY-17464S1

Cilostazol-d4 is deuterium labeled Cilostazol. Cilostazol (OPC 13013) is a potent and selective inhibitor of phosphodiesterase (PDE) 3A, the isoform of PDE 3 in the cardiovascular system, with an IC50 of 0.2 μ M.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Cisplatin

Purity:

Size:

Cinobufagin

(Cinobufagine)

(cis-Platinum; CDDP; cis-Diaminodichloroplatinum)

10 mM × 1 mL, 5 mg, 10 mg

Cisplatin (CDDP) is an antineoplastic chemotherapy agent by cross-linking with DNA and causing DNA damage in cancer cells. Cisplatin activates ferroptosis and induces autophagy.

Cinobufagin, a kind of Chinese materia medica with

antitumor effect, is widely used in clinical practice, especially in anti-liver cancer. IC50

98 90%

Clinical Data: No Development Reported

value: Target: In vitro: Cinobufagin inhibited

proliferation of cancer cells at doses of 0.1, 1, or 10 μM after 2–4 days of culture.

Cat. No.: HY-N0421

Cat. No.: HY-17394

Cat. No.: HY-17508

Purity: >98% Clinical Data: Launched 100 mg, 500 mg

Cisatracurium besylate

(51W89) Cat. No.: HY-13596

Cisatracurium besylate (51W89) is a nondepolarizing neuromuscular blocking agent, antagonizing the action of acetylcholine by inhibiting neuromuscular transmission.

Purity: >98.0% Clinical Data: Launched

10 mM × 1 mL, 25 mg, 50 mg, 100 mg Size:

Citalopram hydrobromide

((±)-Citalopram hydrobromide; Lu 10-171) Cat. No.: HY-B1287

Citalopram hydrobromide is a selective serotonin reuptake inhibitor (SSRI). Citalopram hydrobromide inhibits 5-HT uptake into synaptosomes with an IC_{so} of 1.8 nM. Citalopram hydrobromideinhibits the 5-HT uptake in rabbit blood platelets with an IC₅₀ of 14 nM. Antidepressant effect.



99.66% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg

Clarithromycin

Clarithromycin has a broad spectrum of antimicrobial activity. Clarithromycin inhibits the CYP3A4-catalyzed triazolam alpha-hydroxylation with the IC_{50} (K_i) value of 56 (43) μ M. Clarithromycin significantly inhibits the HERG potassium current.

Purity: >98.0% Clinical Data: Launched

10 mM × 1 mL, 100 mg, 200 mg, 500 mg

Clemastine fumarate

(HS-592 fumarate; Meclastine fumarate) Cat. No.: HY-B0298A

Clemastine (HS-592) fumarate is a selective histamine H1 receptor antagonist. Clemastine fumarate is an antihistamine mainly used for relieving symptoms of allergic reactions primarily by competing with histamine to bind H1 receptors. Anti-inflammatory effects.



Purity: 99.95% Clinical Data: Launched

10 mM \times 1 mL, 100 mg, 200 mg, 500 mg Size:

Clemastine-d5 fumarate

(HS-592-d5 fumarate: Meclastine-d5 fumarate) Cat. No.: HY-B0298AS

Clemastine-d5 (HS-592-d5) fumarate is the deuterium labeled Clemastine fumarate. Clemastine fumarate (HS-592 fumarate) is a selective histamine H1 receptor antagonist with IC₅₀ of 3 nM.

>98% Purity:

Clinical Data: No Development Reported

Size 1 mg, 5 mg

Clematichinenoside AR

Cat. No.: HY-N4232

Clematichinenoside AR is a major active ingredient that could be extracted from the traditional Chinese herb Clematis chinensis and has potent pharmacological effects on various diseases, including atherosclerosis (AS).



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Clinofibrate

(S-8527) Cat. No.: HY-13528

Clinofibrate (S-8527) is a hypelipidemic agent and a HMG-CoA reductase inhibitor.



99.70% Clinical Data: Launched

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

Clionamine B

Clionamine B is an autophagy stimulating aminosteroid isolated from the sponge Cliona celata. Clionamine B strongly stimulates autophagy in human breast cancer MCF-7 cells.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Cat. No.: HY-N10305

Clioquinol (Iodochlorhydroxyquin) is a topical antifungal agent with anticancer activity. Clioquinol acts as an oral antimicrobial agent for the research of diarrhea and skin infections. Antibiotic

98 63% Purity: Clinical Data: Launched

(Iodochlorhydroxyquin)

10 mM × 1 mL, 500 mg, 1 g, 5 g Size:



Cat. No.: HY-14603

Clofarabine

Cat. No.: HY-A0005

Clofarabine, a nucleoside analogue for research of cancer, is a potent inhibitor of ribonucleotide reductase (IC_{so}=65 nM) by binding to the allosteric site on the regulatory subunit.

Purity: 99 09% Clinical Data: Launched

10 mM × 1 mL, 10 mg, 50 mg Size:

Clotrimazole

Clioquinol

Clotrimazole is an imidazole derivative, an antifungal compound and is a CYP (cytochrome P450) inhibitor. Clotrimazole has antibacterial

activity.

Purity: 99 88% Clinical Data: Launched

10 mM × 1 mL, 500 mg, 1 g



Cat. No.: HY-10882

Clotrimazole-d5

Cat. No.: HY-10882S

Clotrimazole-d5 is the deuterium labeled Clotrimazole. Clotrimazole is an imidazole derivative, an antifungal compound and is a CYP (cytochrome P450) inhibitor. Clotrimazole has antibacterial activity.

Purity: >98%

Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg

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 $\alpha 3$ glycine receptors (GlyRs).

99.87% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 200 mg, 500 mg



Cat. No.: HY-16569

Colchicine-d3

Cat. No.: HY-16569S1

Colchicine-d3 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 $\alpha 3$ glycine receptors (GlyRs).

Purity: >98%

Clinical Data: No Development Reported 1 mg, 5 mg, 10 mg Size:

Colchicine-d6

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 $\alpha 3$ glycine receptors (GlyRs).

Purity: >98%

Clinical Data: No Development Reported

Size 1 mg, 10 mg



Cat. No.: HY-16569S

Colistin sulfate

(Polymyxin E Sulfate)

Colistin sulfate is a polypeptide antibiotic which inhibits gram-negative bacteria by binding to lipopolysaccharides and phospholipids in the outer cell membrane of gram-negative bacteria.



Cat. No.: HY-A0089

Purity: ≥96.0% Clinical Data: Launched

Size 10 mM × 1 mL, 100 mg, 500 mg

Cordycepin

(3'-Deoxyadenosine)

Cordycepin (3'-Deoxyadenosine) is a nucleoside derivative and inhibits IL-1 β -induced MMP-1 and MMP-3 expression in rheumatoid arthritis synovial fibroblasts (RASFs) in a dose-dependent manner.

Purity: 98.64% Clinical Data: Phase 2

10 mM × 1 mL, 10 mg, 25 mg, 50 mg, 100 mg



Cat. No.: HY-N0262

www.MedChemExpress.com

Corosolic acid

(Colosolic acid) Cat. No.: HY-N0280

Corosolic acid (Colosolic acid) isolated from the fruit of Cratoegus pinnatifida var. psilosa, was reported to have anticancer activity.



Purity: ≥98.0%

Corynoxine B

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 5 mg, 10 mg

Corynoxine

Corynoxine, a tetracyclic oxindole alkaloid, is isolated from the hooks of Uncaria macrophylla. Corynoxine is a natural autophagy enhancer that promotes the clearance of alpha-synuclein via Akt/mTOR pathway.



Cat. No.: HY-N0901

Purity: 99.58%

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 5 mg, 10 mg

Corynoxine hydrochloride

Corynoxine B is an oxindole alkaloid isolated from Uncaria rhynchophylla (Miq.) Jacks (Gouteng in Chinese); a Beclin-1-dependent autophagy inducer.

Cat. No.: HY-N0901A

Purity: 99.76%

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 5 mq, 10 mq, 25 mq

Corynoxine hydrochloride, a tetracyclic oxindole alkaloid, is isolated from the hooks of Uncaria macrophylla. Corynoxine hydrochloride is a natural autophagy enhancer that promotes the clearance of alpha-synuclein via Akt/mTOR nathway.

TNC H

Cat. No.: HY-N0901B

Purity: >98%

Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg

Crenolanib

(CP-868596) Cat. No.: HY-13223

Crenolanib is a potent and selective inhibitor of wild-type and mutant isoforms of the class III receptor tyrosine kinases FLT3 and PDGFR α/β with K_a s of 0.74 nM and 2.1 nM/3.2 nM, respectively.



Purity: 99.72%

Clinical Data: No Development Reported

Size: $10 \text{ mM} \times 1 \text{ mL}$, 5 mg, 10 mg, 50 mg, 100 mg

Crizotinib

(PF-02341066) Cat. No.: HY-50878

Crizotinib (PF-02341066) is an orally bioavailable, ATP-competitive ALK and **c-Met** inhibitor with $\rm IC_{50}^{5}$ of 20 and 8 nM, respectively.



Purity: 99.97% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg, 500 mg

Crizotinib hydrochloride

(PF-02341066 hydrochloride) Cat. No.: HY-50878A

Crizotinib hydrochloride (PF-02341066 hydrochloride) is an orally bioavailable, selective, and ATP-competitive dual **ALK** and **c-Met** inhibitor with $\rm IC_{50}$ s of 20 and 8 nM, respectively.



Purity: 99.86%
Clinical Data: Launched

Size: $10 \text{ mM} \times 1 \text{ mL}, 10 \text{ mg}, 50 \text{ mg}, 100 \text{ mg}, 200 \text{ mg}, 500 \text{ mg}$

Crizotinib-d5

(PF-02341066-d5) Cat. No.: HY-50878S

Crizotinib-d5 (PF-02341066-d5) is the deuterium labeled Crizotinib. Crizotinib (PF-02341066) is an orally bioavailable, ATP-competitive **ALK** and **c-Met** inhibitor with IC_{50} S of 20 and 8 nM, respectively.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Cryptotanshinone

(Cryptotanshinon; Tanshinone c)

Cryptotanshinone is a natural compound extracted from the root of Salvia miltiorrhiza Bunge that shows antitumor activities. Cryptotanshinone inhibits STAT3 with an IC $_{\rm 50}$ of 4.6 μM .



Cat. No.: HY-N0174

Purity: 98.46%

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 10 mg, 50 mg

Crustecdysone

(20-Hydroxyecdysone)

Crustecdysone (20-Hydroxyecdysone) is a naturally occurring ecdysteroid hormone isolated from Cyanotis arachnoides C.B.Clarke which controls the ecdysis (moulting) and metamorphosis of arthropods, it inhibits caspase activity and induces autophagy via the 20E nuclear...



Cat. No.: HY-N6979

Purity: 99.64%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

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

Cucurbitacin B

Cat. No.: HY-N0416

Cucurbitacin B belongs to a class of highly oxidized tetracyclic triterpenoids; could repress cancer cell progression.

Purity: 99 92%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg

Curcumin

(Diferuloylmethane; Natural Yellow 3; Turmeric yellow) Cat. No.: HY-N0005

Curcumin (Diferuloylmethane), a natural phenolic compound, is a p300/CREB-binding protein-specific inhibitor of acetyltransferase, represses the acetylation of histone/nonhistone proteins and histone acetyltransferase-dependent chromatin transcription.

Purity: ≥96.0% Clinical Data: Phase 4

10 mM × 1 mL, 100 mg, 500 mg Size:

Cucurbitacin E

(α-Elaterin; α-Elaterine)

Cucurbitacin E is a natural compound which from the climbing stem of Cucumic melo L. Cucurbitacin E significantly suppresses the activity of the cyclin B1/CDC2 complex.



Cat. No.: HY-N0417

Purity: 99 92%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg

Curcumin-d6 (Diferuloylmethane-d6; Natural Yellow 3-d6;

Turmeric yellow-d6) Cat. No.: HY-N0005S

Curcumin D6 (Diferuloylmethane D6) is a deuterium labeled Curcumin (Turmeric yellow). Curcumin (Turmeric yellow) is a natural phenolic compound with diverse pharmacologic effects including anti-inflammatory, antioxidant, antiproliferative and antiangiogenic activities.



>98% **Purity:**

Clinical Data: No Development Reported 1 mg, 5 mg, 10 mg

CX546

Cat. No.: HY-12505

CX546 is a first-generation and selective benzamide-type positive AMPAR modulator. CX546 is a prototypical ampakine agent and has antipsychotic effects.

Purity: 99.07%

Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 50 mg

Cyclovirobuxine D

Cyclovirobuxine D (CVB-D) is the main active component of the traditional Chinese medicine Buxus microphylla. Cyclovirobuxine D induces autophagy and attenuates the phosphorylation of Akt and mTOR.



Cat. No.: HY-N0107

≥95.0% Purity:

Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg, 20 mg

Cysteamine hydrochloride (2-Aminoethanethiol hydrochloride;

2-Mercaptoethylamine hydrochloride)

Cysteamine hydrochloride (2-Aminoethanethiol hydrochloride) is an orally active agent for the treatment of nephropathic cystinosis and an antioxidant.

Cat. No.: HY-77591

HCI

≥95.0% Purity: Clinical Data: Launched

10 mM × 1 mL, 500 mg, 5 g Size:

Cysteamine-d4 hydrochloride (2-Aminoethanethiol-d4

hydrochloride; 2-Mercaptoethylamine-d4 hydrochloride) Cat. No.: HY-77591S

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.

$$D$$
 SH D HCI

Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Cytarabine (Cytosine β-D-arabinofuranoside; Cytosine Arabinoside; Ara-C)

Cytarabine, a nucleoside analog, causes S phase cell cycle arrest and inhibits DNA polymerase. Cytarabine inhibits DNA synthesis with an IC_{so} of 16 nM. Cytarabine has antiviral effects against HSV.

Cat. No.: HY-13605

Purity: 99.96% Clinical Data: Launched

10 mM × 1 mL, 100 mg, 500 mg, 1 g

Cytarabine hydrochloride (Cytosine β-D-arabinofuranoside

hydrochloride; Cytosine Arabinoside hydrochloride; ...) Cat. No.: HY-13605A

Cytarabine hydrochloride, a nucleoside analog, causes S phase cell cycle arrest and inhibits DNA polymerase. Cytarabine inhibits DNA synthesis with an IC_{so} of 16 nM. Cytarabine hydrochloride has antiviral effects against HSV.



≥95.0% Clinical Data: Launched

10 mM × 1 mL, 100 mg

Cytarabine-d2

Cat. No.: HY-13605S

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 IC₅₀ of 16 nM. Cytarabine has antiviral effects against

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Cytochalasin E

Cytochalasin E, an epoxide containing Aspergillus-derived fungal metabolite, inhibits angiogenesis and tumor growth. Cytochalasin E is a potent actin depolymerization agent, and it binds and caps the barbed end of actin filaments to prevent actin elongation.

Purity: >98%

Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg



Cat. No.: HY-N6772

D-Glutamine

Cat. No.: HY-100587

D-Glutamine is a cell-permeable D type stereoisomer of Glutamine.

Purity: > 98.0%

Clinical Data: No Development Reported 10 mM × 1 mL, 50 mg, 100 mg

D4476

(Casein Kinase I Inhibitor)

D4476 is a potent, selective and cell-permeable inhibitor of casein kinase 1(CK1) with an IC50 value of 0.3 μM in vitro.

Cat. No.: HY-10324

Purity: 99 51%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg

Dacinostat

(NVP-LAQ824; LAQ824) Cat. No.: HY-13606

Dacinostat is a potent HDAC inhibitor, with an IC₅₀ of 32 nM; Dacinostat also inhibits HDAC1 with an IC_{50} of 9 nM, and used in cancer research.

98.45% Purity:

Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 50 mg

Dactolisib

(BEZ235; NVP-BEZ235)

Dactolisib (BEZ235) is an orally active and dual pan-class I PI3K and mTOR kinase inhibitor with IC_{so}s of 4 nM/5 nM/7 nM/75 nM, and 20.7 nM for $p110\alpha/p110\gamma/p110\delta/p110\beta$ and mTOR, respectively. Dactolisib (BEZ235) inhibits both mTORC1 and mTORC2.

Purity: 99 94% Clinical Data: Phase 3

Size 50 mg, 100 mg, 200 mg, 500 mg



Cat. No.: HY-50673

Dactolisib Tosylate

(BEZ235 Tosylate; NVP-BEZ 235 Tosylate) Cat. No.: HY-15174

Dactolisib Tosylate (BEZ235 Tosylate) is a dual PI3K and mTOR kinase inhibitor with IC₅₀ values of 4, 75, 7, 5 nM for PI3K α , β , γ , δ , respectively. Dactolisib Tosylate (BEZ235 Tosylate) inhibits mTORC1 and mTORC2.

99.88% Purity: Clinical Data: Phase 3

10 mM \times 1 mL, 50 mg, 100 mg, 200 mg, 500 mg Size

Danshensu

(Dan shen suan A; Salvianic acid A)

Danshensu, an active ingredient of Salvia miltiorrhiza, shows wide cardiovascular benefit by activating Nrf2 signaling pathway.

Cat. No.: HY-N1913

≥98.0% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 10 mg, 50 mg, 100 mg

Danshensu sodium salt

(Sodium Danshensu; (±)-DanShenSu sodium sal) Cat. No.: HY-N0106

Danshensu (sodium salt) is odium salt of danshensu from the widely used Chinese herb Danshen. It can inhibited phenylephrine- and CaCl2-induced vasoconstriction in Ca2+-free medium. In vitro: Sodium danshensu showed a biphasic effects on vessel tension.



Purity: ≥98.0%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg

Dansylcadaverine

(Monodansyl cadaverine)

Dansylcadaverine (Monodansyl cadaverine) is an autofluorescent compound used for the labeling of autophagic vacuoles. Dansylcadaverine, a high affinity substrate of transglutaminases, can block the receptor-mediated endocytosis of many ligands.



Cat. No.: HY-D1027

98.62%

Clinical Data: No Development Reported 10 mM × 1 mL, 50 mg

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Danthron

(Dantron; Chrysazin; 1,8-Dihydroxyanthraquinone)

Danthron is a natural product extracted from the traditional Chinese medicine rhubarb. Danthron functions in regulating glucose and lipid metabolism by activating AMPK.

Cat. No.: HY-B0923

98 70% Purity: Clinical Data: Launched

Size: $10 \text{ mM} \times 1 \text{ mL}, 100 \text{ mg}$

Danthron-d6

(Dantron-d6; Chrysazin-d6; 1,8-Dihydroxyanthraquinone-d6) Cat. No.: HY-B0923S

Danthron-d6 (Dantron-d6) is the deuterium labeled Danthron, Danthron is a natural product extracted from the traditional Chinese medicine rhubarb. Danthron functions in regulating glucose and lipid metabolism by activating AMPK.

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 10 mg

Dantrolene sodium hemiheptahydrate

(Dantrolene sodium hydrate)

Dantrolene sodium hemiheptahydrate is a skeletal muscle relaxant which acts by blocking muscle contraction beyond the neuromuscular junction. Dantrolene sodium hemiheptahydrate is a inhibitor of calcium channel proteins, inhibiting the release of Ca2+ from the sarcoplasm.

Cat. No.: HY-12542A

Purity: > 98.0% Clinical Data: Launched

10 mM × 1 mL, 100 mg, 200 mg, 500 mg Size:

Danusertib

(PHA-739358) Cat. No.: HY-10179

Danusertib is a pyrrolo-pyrazole and aurora kinase inhibitor with IC₅₀ of 13, 79, and 61 nM for Aurora A, B, and C, respectively.



Purity: 99 44% Clinical Data: Phase 2

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Daphnetin

(7,8-Dihydroxycoumarin) Cat. No.: HY-N0281

Daphnetin (7,8-dihydroxycoumarin), one coumarin derivative isolated from plants of the Genus Daphne, is a protein kinase inhibitor, with IC₅₀s of 7.67 μM, 9.33 μM and 25.01 μM for EGFR, PKA and PKC in vitro, respectively.



Purity: 99 21% Clinical Data: Launched

Size: $10 \text{ mM} \times 1 \text{ mL}$, 10 mg, 50 mg, 100 mg

Dapivirine

(TMC120; R147681)

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.



Cat. No.: HY-14266

99 90% Purity: Clinical Data: Phase 3

Size $10~\text{mM}\times1~\text{mL},\,5~\text{mg},\,10~\text{mg},\,50~\text{mg},\,100~\text{mg}$

Dapivirine-d11

(TMC120-d11; R147681-d11) Cat. No.: HY-14266S

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: >98%

Clinical Data: No Development Reported

Size 1 ma, 10 ma

DAPT

(GSI-IX) Cat. No.: HY-13027

DAPT (GSI-IX) is a potent and orally active γ -secretase inhibitor with IC₅₀s 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.



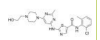
Purity: 99.93%

Clinical Data: No Development Reported 10 mM \times 1 mL, 5 mg, 10 mg, 50 mg

Dasatinib

(BMS-354825) Cat. No.: HY-10181

Dasatinib (BMS-354825) is a highly potent, ATP competitive, orally active dual Src/Bcr-Abl inhibitor with potent antitumor activity. The Kis are 16 pM and 30 pM for Src and Bcr-Abl, respectively.



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 (BMS-354825) hydrochloride is a highly

potent, ATP competitive, orally active dual Src/Bcr-Abl inhibitor with potent antitumor activity. The $\mathbf{K}_{\mathrm{i}}\mathrm{s}$ are 16 pM and 30 pM for Src and Bcr-Abl, respectively.



Purity: 98.86% Clinical Data: Launched

10 mM × 1 mL, 100 mg, 200 mg, 500 mg

Dasatinib monohydrate

(BMS-354825 monohydrate) Cat. No.: HY-10181B

Dasatinib (BMS-354825) monohydrate is a highly potent, ATP competitive, orally active dual Src/Bcr-Abl inhibitor with potent antitumor activity. The K_is are 16 pM and 30 pM for Src and Bcr-Abl, respectively.

>98% Purity: Clinical Data: Launched Size: 1 mg, 5 mg

Daunorubicin

(Daunomycin; RP 13057; Rubidomycin) Cat. No.: HY-13062A

Daunorubicin (Daunomycin; RP 13057; Rubidomycin) is a topoisomerase II inhibitor with potent antineoplastic activities. Daunorubicin (Daunomycin; RP 13057; Rubidomycin) inhibits DNA and RNA synthesis in sensitive and resistant Ehrlich ascites tumor cells.



Purity: >98% Clinical Data: Launched 5 mg, 10 mg, 25 mg

Daurisoline

((R,R)-Daurisoline) Cat. No.: HY-N0221

Daurisoline is a hERG inhibitor and also an autophagy blocker.



Purity: 99.65%

DC-LC3in-D5

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg

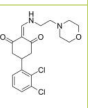
Cat. No.: HY-141882

DC-LC3in-D5 acts as an autophagy inhibitor by attenuating LC3B lipidation. DC-LC3in-D5 binds with LC3B. DC-LC3in-D5 disrupts the LC3B-LBP2 interaction with an IC₅₀ of 200 nM. DC-LC3in-D5 may contribute to anti-HCV or combination treatments in cancer through inhibiting autophagy.

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg



Deferoxamine mesylate

(Desferrioxamine B mesylate; DFOM) Cat. No.: HY-B0988

Deferoxamine mesylate is an iron chelator that binds free iron in a stable complex, preventing it from engaging in chemical reactions.

Purity: 99.86% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg

Dasatinib-d8

(BMS-354825-d8) Cat. No.: HY-10181S

Dasatinib D8 is a deuterium labeled Dasatinib. Dasatinib is a dual Bcr-Abl and Src family tyrosine kinase inhibitor.



>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Daunorubicin hydrochloride (Daunomycin hydrochloride; RP

13057 hydrochloride; Rubidomycin hydrochloride) Cat. No.: HY-13062

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

10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg, 500 mg Size:

DBeQ

(JRF 12) Cat. No.: HY-15945

DBeQ is a selective, potent, reversible, and ATP-competitive p97 inhibitor, with an IC₅₀ value of 1.5 μM and 1.6 μM for p97(wt) and p97(C522A), respectively; DBeQ also inhibits Vps4 with an IC_{50} of 11.5 μM .



99.68% Purity:

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

DC661

Cat. No.: HY-111621

DC661 is a potent palmitoyl-protein thioesterase 1 (PPT1) inhibitor, inhibits autophagy, and acts as an anti-lysosomal agent. Anti-cancer activity.



Purity: ≥95.0%

Clinical Data: No Development Reported

10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Degrasyn

(WP1130) Cat. No.: HY-13264

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.



99.70% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Deguelin

((-)-Deguelin; (-)-cis-Deguelin)

Dequelin, a naturally occurring rotenoid, acts as a chemopreventive agent by blocking multiple pathways like PI3K-Akt, IKK-NF-κB, and MAPK-mTOR-survivin-mediated apoptosis.

Cat. No.: HY-13425

Purity: 99 29%

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

Dehydrocorydaline

(13-Methylpalmatine)

Dehydrocorydaline (13-Methylpalmatine) is an alkaloid that regulates protein expression of Bax, Bcl-2; activates caspase-7, caspase-8, and inactivates PARP. Dehydrocorydaline elevates p38 MAPK activation. Anti-inflammatory and anti-cancer activities.



Cat. No.: HY-N0674

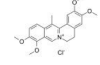
Purity: 99.01%

Clinical Data: No Development Reported 10 mM × 1 mL, 5 mg, 10 mg Size:

Dehydrocorydaline chloride

(13-Methylpalmatine chloride)

Dehydrocorydaline chloride (13-Methylpalmatine chloride) is an alkaloid that regulates protein expression of Bax, Bcl-2; activates caspase-7, caspase-8, and inactivates PARP. Dehydrocorydaline chloride elevates p38 MAPK activation.



Cat. No.: HY-N0674A

Purity:

Clinical Data: No Development Reported 10 mM × 1 mL, 5 mg, 10 mg

Dehydrocorydaline nitrate

(13-Methylpalmatine nitrate)

Dehydrocorydaline nitrate (13-Methylpalmatine nitrate) is an alkaloid. Dehydrocorydaline regulates protein expression of Bax, Bcl-2; activates caspase-7, caspase-8, and inactivates PARP. Dehydrocorydaline nitrate elevates p38 MAPK activation.



Clinical Data: No Development Reported

5 mg, 10 mg



Cat. No.: HY-N4238

Dehydropachymic acid

Cat. No.: HY-N2991

Dehydropachymic acid is one of the major triterpenes isolated from Poria cocos. Dehydropachymic acid is more effective in autophagy-lysosome pathway (ALP) impaired cells rather than normal cells.



Purity: 99 94%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Demethoxycurcumin

(Curcumin II; Desmethoxycurcumin; Monodemethoxycurcumin) Cat. No.: HY-N0006

Demethoxycurcumin(Curcumin II) is a major active curcuminoid; possess anti-inflammatory properties; also exert cytotoxic effects in human cancer cells via induction of apoptosis. IC50 value: Target: in vitro: DMC significantly decreased NO secretion by 35-41% in our inflamed cell model.



Purity: ≥99.0%

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

Demethoxycurcumin-d7 (Curcumin II-d7; Desmethoxycurcumin-d7; Cat. No.: HY-N0006S

Monodemethoxycurcumin-d7)

Demethoxycurcumin-d7 (Curcumin II-d7) is the deuterium labeled Demethoxycurcumin. Demethoxycurcumin(Curcumin II), a major active curcuminoid, possess anti-inflammatory properties; also exert cytotoxic effects in human cancer cells via induction of apoptosis.



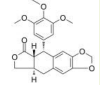
Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Deoxypodophyllotoxin

Deoxypodophyllotoxin (DPT), a derivative of podophyllotoxin, is a lignan with potent antimitotic, anti-inflammatory and antiviral properties isolated from rhizomes of Sinopodophullumhexandrum (Berberidaceae).



Cat. No.: HY-N2500

99.86% Purity:

Clinical Data: No Development Reported

5 mg, 10 mg

Desethyl Amiodarone-d4 hydrochloride

Cat. No.: HY-130353S

Desethyl Amiodarone-d4 hydrochloride is the deuterium labeled Desethylamiodarone hydrochloride. Desethylamiodarone hydrochloride (N-desethylamiodarone hydrochloride) is a major active metabolite of Amiodarone.



Purity: >98%

Clinical Data: No Development Reported

Size 1 mg, 10 mg

Desethyl chloroquine

Cat. No.: HY-135811

Desethyl chloroquine is a major desethyl metabolite of Chloroquine. Chloroquine diphosphate is an inhibitor of autophagy and toll-like receptors (TLRs). Desethyl chloroquine possesses antiplasmodic activity.



Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg

Desethyl chloroquine diphosphate

Cat. No.: HY-135811A

Desethyl chloroquine diphosphate is a major desethyl metabolite of Chloroquine. Chloroquine diphosphate is an inhibitor of **autophagy** and **toll-like receptors (TLRs)**. Desethyl chloroquine diphosphate possesses antiplasmodic activity.

Purity: 99.44%

Clinical Data: No Development Reported
Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Desethyl chloroquine-d4

Desethyl chloroquine-d4 is the deuterium labeled Desethyl chloroquine. Desethyl chloroquine is a major desethyl metabolite of Chloroquine. Chloroquine diphosphate is an inhibitor of autophagy and toll-like receptors (TLRs).



Cat. No.: HY-135811S

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 10 mg

Desethyl chloroquine-d5

Cat. No.: HY-135811S1

Desethyl chloroquine-d5 is deuterium labeled Desethyl chloroquine. Desethyl chloroquine is a major desethyl metabolite of Chloroquine. Chloroquine diphosphate is an inhibitor of autophagy and toll-like receptors (TLRs). Desethyl chloroquine possesses antiplasmodic activity.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Desethylamiodarone hydrochloride (N-desethylamiodarone

hydrochloride; LB 33020 hydrochloride)

Desethylamiodarone hydrochloride (N-desethylamiodarone hydrochloride) is a major active metabolite of Amiodarone.
Desethylamiodarone hydrochloride is formed by CYP3A isoenzymes.



Cat. No.: HY-130353

Purity: ≥98.0%

Clinical Data: No Development Reported

Size: 5 mg, 10 mg

Desfluoro-ezetimibe

Cat. No.: HY-136059

Desfluoro-ezetimibe is a desfluoro impurity of Ezetimibe. Ezetimibe is a potent, metabolically stable cholesterol absorption inhibitor. Ezetimibe is a Niemann-Pick C1-like1 (NPC1L1) inhibitor, and is a potent Nrf2 activator.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Desmethyl Naproxen-d3

Cat. No.: HY-132405S

Desmethyl Naproxen-d3 is deuterium labeled Desmethyl Naproxen. Desmethyl Naproxen is the metabolite of anti-inflammatory agent Naproxen.

Purity: ≥98.0%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Dexamethasone

(Hexadecadrol; Prednisolone F)

Dexamethasone (Hexadecadrol) is a **glucocorticoid receptor** agonist. Dexamethasone also significantly decreases CD11b, CD18, and CD62L expression on neutrophils, and CD11b and CD18 expression on monocytes.



Cat. No.: HY-14648

Purity: 99.86% Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g

Dexamethasone acetate

(Dexamethasone 21-acetate; Hexadecadrol acetate) Cat. No.: HY-14648A

Dexamethasone acetate (Dexamethasone 21-acetate) is a **glucocorticoid receptor** agonist.

Dexamethasone acetate has the potential for ophthalmic infections treatment.



Purity: 98.24% Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g

Dexamethasone-4,6 α ,21,21-d4

Cat. No.: HY-14648S3

Dexamethasone- $4,6\alpha,21,21$ -d4 is the deuterium labeled Dexamethasone- $4,6\alpha,21,21$. Dexamethasone (Hexadecadrol) is a glucocorticoid receptor agonist.



Purity: > 98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Dexamethasone-d4

(Hexadecadrol-d4; Prednisolone F-d4)

Dexamethasone-d4 is deuterium labeled Dexamethasone. Dexamethasone (Hexadecadrol) is a gluccocrticoid receptor agonist. Dexamethasone also significantly decreases CD11b, CD18, and CD62L expression on neutrophils, and CD11b and CD18 expression on monocytes.



Cat. No.: HY-14648S2

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

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Dexamethasone-d5

(Hexadecadrol-d5; Prednisolone F-d5)

Dexamethasone-d5 (Hexadecadrol-d5) is the deuterium labeled Dexamethasone. Dexamethasone (Hexadecadrol) is a glucocorticoid receptor agonist.



Cat. No.: HY-14648S

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 10 mg

Diazoxide

(Sch-6783; SRG-95213) Cat. No.: HY-B1140

Diazoxide (Sch-6783) is an ATP-sensitive potassium channel activator, has the potential for hyperinsulinism treatment.

Purity: 99 99% Clinical Data: Launched

10 mM × 1 mL, 100 mg

Diazoxide-d3

Purity:

Size:

(Sch-6783-d3; SRG-95213-d3)

CD18 expression on monocytes.

>98% Clinical Data: No Development Reported

1 mg, 5 mg

Dexamethasone-d5-1

(Hexadecadrol-d5-1; Prednisolone F-d5-1)

Dexamethasone-d5-1 is deuterium labeled

Dexamethasone. Dexamethasone (Hexadecadrol) is a

glucocorticoid receptor agonist. Dexamethasone also significantly decreases CD11b, CD18, and

CD62L expression on neutrophils, and CD11b and

Diazoxide-d3 is deuterium labeled Diazoxide. Diazoxide (Sch-6783) is an ATP-sensitive potassium channel activator, has the potential for hyperinsulinism treatment.

Cat. No.: HY-B1140S

Cat. No.: HY-14648S1

Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg

Dienogest

(STS 557) Cat. No.: HY-B0084

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).



99.83% Purity: Clinical Data: Launched

Size: $10 \text{ mM} \times 1 \text{ mL}$, 10 mg, 50 mg, 100 mg

Dienogest-d4

(STS 557-d4) Cat. No.: HY-B0084S

Dienogest-d4 is deuterium labeled Dienogest.



>98% Purity:

Clinical Data: No Development Reported

Size 1 mg, 5 mg

Dienogest-d5

(STS 557-d5) Cat. No.: HY-B0084S1

Dienogest-d5 is deuterium labeled Dienogest.



>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Dienogest-d6

(STS 557-d6) Cat. No.: HY-B0084S2

Dienogest-d6 is deuterium labeled Dienogest.

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Dihydroartemisinin

(Dihydroqinghaosu; β-Dihydroartemisinin; Artenimol) Cat. No.: HY-N0176

Dihydroartemisinin is a potent anti-malaria agent.



Purity: ≥98.0% Clinical Data: Launched

Size: 10 mM × 1 mL, 50 mg, 100 mg, 200 mg, 500 mg

Dihydroartemisinin-d3 (Dihydroqinghaosu-d3;

β-Dihydroartemisinin-d3; Artenimol-d3)

Dihydroartemisinin-d3 (Dihydroginghaosu-d3) is the deuterium labeled Dihydroartemisinin. Dihydroartemisinin is a potent anti-malaria agent.



Cat. No.: HY-N0176S

>98%

Clinical Data: No Development Reported

1 mg, 5 mg

Dihydromyricetin

(Ampelopsin; Ampeloptin) Cat. No.: HY-N0112

Dihydromyricetin is a potent inhibitor with an IC_{50} of 48 μ M on **dihydropyrimidinase**. Dihydromyricetin can activate autophagy through inhibiting **mTOR** signaling. Dihydromyricetin suppresses the formation of mTOR complexes (mTORC1/2).

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Purity: 99.79% Clinical Data: Phase 2

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Dilmapimod

(SB-681323; GW 681323)

Dilmapimod (SB-681323) is a potent **p38 MAPK** inhibitor that potentially suppresses inflammation in chronic obstructive pulmonary disease.



Cat. No.: HY-10404

Purity: 99.56% Clinical Data: Phase 2

Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Dimethyl fumarate

Cat. No.: HY-17363

Dimethyl fumarate (DMF) is an orally active and brain-penetrant **Nrf2** activator and induces upregulation of antioxidant gene expression.

Purity: 99.88% Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 5 g

Dinoprost

(Prostaglandin F2α; PGF2α)

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).

HQ OH

Cat. No.: HY-12956S

Cat. No.: HY-12956

Purity: 99.06% Clinical Data: Launched

Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Dinoprost tromethamine salt (Prostaglandin F2α tromethamine salt; PGF2α THAM) Cat. No.: HY-12956A

Dinoprost tromethamine salt (Prostaglandin F2α tromethamine salt) is an orally active, potent prostaglandin F (PGF) receptor (FP receptor)

agonist.

HO OH HO NF:

Purity: ≥98.0% Clinical Data: Launched

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Dinoprost-d4

(Prostaglandin F2a-d4; PGF2α-d4)

Dinoprost-d4 (Prostaglandin F2a-d4) is the deuterium labeled Dinoprost. Dinoprost (Prostaglandin F2α) is an orally active, potent prostaglandin F (PGF) receptor (FP receptor) agonist.

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Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Dinoprost-d9

(Prostaglandin F2a-d9; PGF2α-d9) Cat. No.: HY-12956S1

Dinoprost-d9 (Prostaglandin F2a-d9) is the deuterium labeled Dinoprost. Dinoprost (Prostaglandin F2α) is an orally active, potent **prostaglandin F (PGF) receptor (FP receptor)** agonist.

HO OHD DD D

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Dioscin

(Collettiside III; CCRIS 4123)

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.

Cat. No.: HY-N0124

Purity: 99.76% Clinical Data: Launched

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg

Diosgenin glucoside

Cat. No.: HY-N0730

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 .

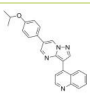


Purity: 99.28%

Clinical Data: No Development Reported Size: 5 mg, 10 mg, 20 mg

DMH-1

DMH-1 is a potent and selective **BMP** inhibitor with IC_{so} s of 27/107.9/<5/47.6 nM for ALK1/ALK2/ALK3/ALK6, respectively.



Cat. No.: HY-12273

Purity: 99.81%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg

DMOG

(Dimethyloxallyl Glycine) Cat. No.: HY-15893

DMOG (Dimethyloxallyl Glycine) is a cell permeable and competitive inhibitor of HIF-PH, which results in HIF-1 α stabilisation and accmulation in vitro and in vivo. DMOG is an α -ketoglutarate analogue and inhibits α -KG-dependent hydroxylases.

Purity: 98.70%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 50 mg, 100 mg, 200 mg, 500 mg

Dorsomorphin

(Compound C; BML-275) Cat. No.: HY-13418A

Dorsomorphin (Compound C) is a selective and ATP-competitive AMPK inhibitor (K_i=109 nM in the absence of AMP). Dorsomorphin (BML-275) selectively inhibits BMP type I receptors ALK2, ALK3, and ALK6. Dorsomorphin induces autophagy



Purity: 99.91%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Doxazosin mesylate

(UK 33274 mesylate) Cat. No.: HY-B0098A

Doxazosin mesylate (UK 33274) is a quinazoline-derivative that selectively antagonizes postsynaptic α1-adrenergic receptors.



Purity: 99.72% Clinical Data: Launched

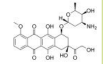
Size: 10 mM × 1 mL, 500 mg, 1 g

.: HY-B0098A (Hydroxyd

(Hydroxydaunorubicin) Cat. No.: HY-15142A

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Doxorubicin (Hydroxydaunorubicin), a cytotoxic anthracycline antibiotic, is an anti-cancer chemotherapy agent. Doxorubicin inhibits $topoisomerase \: I\!I \: with an \: IC_{so} \: of \: 2.67 \: \mu M, \: thus stopping DNA replication.$



Cat. No.: HY-10320

Cat. No.: HY-13418

Purity: >98%
Clinical Data: Launched

Doramapimod

Doramapimod (BIRB 796) is an orally active, highly potent p38 MAPK inhibitor, which has an ${\rm IC_{50}}$

for p38 α =38 nM, for p38 β =65 nM, for p38 γ =200 nM,

and for p38 δ =520 nM. Doramapimod has picomolar

(Compound C dihydrochloride; BML-275 dihydrochloride)

10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg

affinity for p38 kinase ($\rm K_a$ =0.1 nM). Doramapimod also inhibits B-Raf with an IC $_{\rm so}$ of 83 nM.

99.88%

Dorsomorphin dihydrochloride

Dorsomorphin dihydrochloride (BML-275

inhibitor, with a K, of 109 nM.

99 91%

Clinical Data: No Development Reported

potent, selective and ATP-competitive AMPK

dihydrochloride; Compound C dihydrochloride) is a

Clinical Data: Phase 2

(BIRB 796)

Purity:

Size:

Purity:

Doxorubicin

Size: 5 mg, 10 mg, 25 mg

Doxorubicin hydrochloride

(Hydroxydaunorubicin hydrochloride) Cat. No.: HY-15142

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_{50} S of 0.8 μ M and 2.67 μ M, respectively.



Purity: 99.47%
Clinical Data: Launched

Size: 10 mM × 1 mL, 50 mg, 100 mg, 200 mg, 500 mg, 1 g

DPN

(Diarylpropionitrile) Cat. No.: HY-12452

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.



Purity: 99.66%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 500 mg

Dronedarone

(SR 33589) Cat. No.: HY-A0016

Dronedarone (SR 33589), a derivative of amiodarone (HY-14187), is a class III antiarrhythmic agent for the study of atrial fibrillation (AF) and atrial flutter.



Purity: 99.81% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg

Dronedarone D6 hydrochloride

Cat. No.: HY-A0016S

Dronedarone D6 hydrochloride is the deuterium labeled Dronedarone. Dronedarone hydrochloride, a derivative of Amiodarone (HY-14187), is a class III antiarrhythmic agent for the study of atrial fibrillation (AF) and atrial flutter.



urity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Dronedarone Hydrochloride

Cat. No.: HY-75839

Dronedarone Hydrochloride is a non-iodinated amiodarone derivative that inhibits Na+, K+ and Ca2+ currents.

Purity: 99 93% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg

Dynasore

Cat. No.: HY-15304

Dynasore is a cell-permeable dynamin inhibitor with an IC_{so} of 15 μM.



98 70% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 10 mg, 50 mg

F-64

(Proteinase inhibitor E 64) Cat. No.: HY-15282

E-64 (Proteinase inhibitor E 64) is a potent irreversible inhibitor against general cysteine proteases with IC₅₀ of 9 nM for papain.

Purity: 99 96%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg

EACC

Cat. No.: HY-129111

EACC is a reversible autophagy inhibitor, which can block autophagic flux. EACC selectively inhibits the translocation of autophagosome-specific SNARE Stx17 thereby blocking autophagosome-lysosome fusion.

Purity: 99.25%

Clinical Data:

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

EAD1

Cat. No.: HY-123056

EAD1 is a potent autophagy inhibitor with antiproliferative activity in lung and pancreatic cancer cells. EAD1 also induces apoptosis.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Efavirenz

(DMP 266; EFV; L-743726)

Efavirenz (DMP 266) is a potent inhibitor of the wild-type HIV-1 reverse transcriptase with a K, of 2.93 nM and exhibits an IC₉₅ of 1.5 nM for the inhibition of HIV-1 replicative spread in cell culture.

99.84% Purity: Clinical Data: Launched

eIF4A3-IN-2

 $10 \text{ mM} \times 1 \text{ mL}$, 5 mg, 10 mg, 50 mgSize



Cat. No.: HY-10572

Efavirenz-d5

Cat. No.: HY-10572S

Efavirenz-d5 (DMP 266-d5) is the deuterium labeled Efavirenz. Efavirenz (DMP 266) is a potent inhibitor of the wild-type HIV-1 reverse transcriptase with a K_i of 2.93 nM and exhibits an IC_{95} of 1.5 nM for the inhibition of HIV-1 replicative spread in cell culture.

Purity: >98% Clinical Data:

Size: 500 μg, 5 mg

eIF4A3-IN-2 is a highly selective and noncompetitive eukaryotic initiation factor 4A-3 (eIF4A3) inhibitor with an IC₅₀ of 110 nM.

Purity: 99.77%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg Size:



Cat. No.: HY-101785

Elaiophylin

(Azalomycin B; Gopalamicin; Efomycin E) Cat. No.: HY-15184

Elaiophylin (Azalomycin B; Gopalamicin; Efomycin E) is an autophagy inhibitor, exerts antitumor activity as a single agent in ovarian cancer cells.



Purity: 96.20%

Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg, 25 mg

Elexacaftor

(VX-445) Cat. No.: HY-111772

Elexacaftor (VX-445, Compound 1) is a modulator of cystic fibrosis transmembrane conductance regulator (CFTR). Elexacaftor (VX-445, Compound 1) facilitates the processing and trafficking of CFTR to increase the amount of CFTR at the cell surface.

Purity: 99.50% Clinical Data: Launched

10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 50 mg

Embelin

(Embelic acid; Emberine; NSC 91874) Cat. No.: HY-17473

Embelin (Embelic acid), a potent, nonpeptidic XIAP inhibitor (IC_{so} =4.1 μ M), inhibits cell growth, induces apoptosis, and activates caspase-9 in prostate cancer cells with high levels of XIAP.

Purity: 98 75%

Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 50 mg

Emodin

(Frangula emodin)

Emodin (Frangula emodin), an anthraquinone derivative, is an anti-SARS-CoV compound, Emodin blocks the SARS coronavirus spike protein and angiotensin-converting enzyme 2 (ACE2) interaction. Emodin inhibits casein kinase-2 (CK2). Anti-inflammatory and anticancer effects.



Cat. No.: HY-14393

99 39% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 50 mg, 100 mg, 200 mg

Emodin-d4

(Frangula emodin-d4) Cat. No.: HY-14393S

Emodin-d4 (Frangula emodin-d4) is the deuterium labeled Emodin. Emodin (Frangula emodin), an anthraquinone derivative, is an anti-SARS-CoV compound. Emodin blocks the SARS coronavirus spike protein and angiotensin-converting enzyme 2 (ACE2) interaction.

Purity: >98%

Clinical Data: No Development Reported

Size 1 mg, 10 mg

EN6

EN6 is a small-molecule in vivo activator of

autophagy that covalently targets cysteine 277 in the ATP6V1A subunit of the lysosomal the vacuolar

H+ ATPase (v-ATPase).

Cat. No.: HY-128892

Purity: 99.16%

Clinical Data:

Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 50 mg, 100 mg

Enalaprilat dihydrate

(MK-422) Cat. No.: HY-B0231

Enalaprilat dihydrate (MK-422) is an angiotensin-converting enzyme (ACE) inhibitor with IC₅₀ of 1.94 nM.

Purity: 99.68% Clinical Data: Launched

 $10 \text{ mM} \times 1 \text{ mL}$, 50 mg, 100 mg, 500 mgSize:

Entinostat

(MS-275; SNDX-275) Cat. No.: HY-12163

Entinostat is an oral and selective class I HDAC inhibitor, with IC_{50} s of 243 nM, 453 nM, and 248 nM for HDAC1, HDAC2, and HDAC3, respectively.



99.65% Purity: Clinical Data: Phase 3

 $10~\text{mM}\times1~\text{mL},\,10~\text{mg},\,50~\text{mg},\,100~\text{mg},\,200~\text{mg}$ Size:

Entrectinib

(NMS-E628; RXDX-101) Cat. No.: HY-12678

Entrectinib (NMS-E628) is a potent, orally available, and CNS-active pan-Trk, ROS1, and ALK inhibitor. Entrectinib inhibits TrkA, TrkB, TrkC, ROS1 and ALK with IC₅₀ values of 1, 3, 5, 12 and 7 nM, respectively. Antitumor activity.



99.32% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Enzalutamide

(MDV3100) Cat. No.: HY-70002

Enzalutamide (MDV3100) is an androgen receptor (AR) antagonist with an IC₅₀ of 36 nM in LNCaP prostate cells. Enzalutamide is an autophagy activator.

99.96% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

Enzalutamide-d3

(MDV3100-d3) Cat. No.: HY-70002S

Enzalutamide D3 is a deuterium labeled Enzalutamide (MDV3100). Enzalutamide is an androgen receptor (AR) antagonist with an IC₅₀ of 36 nM in LNCaP prostate cells.



>98% Purity:

Clinical Data: No Development Reported

1 mg, 5 mg Size:

Enzastaurin

(LY317615)

Enzastaurin (LY317615) is a potent and selective PKCβ inhibitor with an IC_{so} of 6 nM, showing 6to 20-fold selectivity over PKCα, PKCγ and PKCε.



Cat. No.: HY-10342

99.92% **Purity:** Clinical Data: Phase 3

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

Epi Lovastatin-d3

Cat. No.: HY-N0504S

Epi Lovastatin-d3 is the deuterium labeled Lovastatin, Lovastatin is a cell-permeable HMG-CoA reductase inhibitor used to lower cholesterol.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 10 mg

Eprenetapopt

(APR-246; PRIMA-1Met) Cat. No.: HY-19980

Eprenetapopt (APR-246) is a first-in-class, small molecule that restores wild-type p53 functions in TP53-mutant cells. Eprenetapopt triggers apoptosis in tumor cells.



Purity: > 98.0% Clinical Data: Phase 3

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Erlotinib

Purity:

Size:

Episilvestrol

(CP-358774; NSC 718781; OSI-774)

99.86%

Clinical Data: No Development Reported

 $10 \text{ mM} \times 1 \text{ mL}, 1 \text{ mg}$

Episilvestrol is a derivative of silvestrol, isolated from the fruits and twigs of Aglaia

silvestris, and is a specific eIF4A-targeting

translation inhibitor, with antitumor activity.

Erlotinib (CP-358774) is a directly acting EGFR tyrosine kinase inhibitor, with an IC₅₀ of 2 nM for human EGFR. Erlotinib reduces EGFR autophosphorylation in intact tumor cells with an IC₅₀ of 20 nM. Erlotinib is used for the treatment of non-small cell lung cancer.



10 mM × 1 mL, 100 mg, 500 mg



Cat. No.: HY-50896

Cat. No.: HY-15359

Erlotinib Hydrochloride (CP-358774 hydrochloride; NSC 718781

hydrochloride; OSI-774 hydrochloride) Cat. No.: HY-12008

Erlotinib Hydrochloride (CP-358774 Hydrochloride) inhibits purified EGFR kinase with an IC₅₀ of 2 nM.



Purity: 99 99% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg

Erlotinib mesylate (CP-358774 mesylate; NSC 718781 mesylate;

OSI-774 mesylate) Cat. No.: HY-12008A

Erlotinib mesylate (CP-358774 mesylate) inhibits purified EGFR kinase with an IC₅₀ of 2 nM.



>98% Purity: Clinical Data: Launched Size 1 mg, 5 mg

Erlotinib-13C6

(CP-358774-13C6; NSC 718781-13C6; OSI-774-13C6) Cat. No.: HY-50896S1

Erlotinib-13C6 (CP-358774-13C6) is a 13C-labeled Erlotinib. Erlotinib is a directly acting EGFR tyrosine kinase inhibitor, with an IC₅₀ of 2 nM for human EGFR.



Cat. No.: HY-B0957

>98% Purity:

Clinical Data: No Development Reported

Size:

Erlotinib-d6 hydrochloride (CP-358774-d6 hydrochloride; NSC

718781-d6 hydrochloride; OSI-774-d6 hydrochloride) Cat. No.: HY-12008S

Erlotinib D6 hydrochloride (CP-358774 D6 hydrochloride) a deuterium labeled Erlotinib Hydrochloride. Erlotinib Hydrochloride inhibits purified EGFR kinase with an IC₅₀ of 2 nM.



98.13% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Erythromycin Ethylsuccinate

(Erythromycin ethyl succinate; EES)

Erythromycin Ethylsuccinate is an antibiotic useful for the treatment of a number of bacterial infections, has an antimicrobial spectrum similar to or slightly wider than that of penicillin. Erythromycin Ethylsuccinate has antiviral activity against HIV-1.

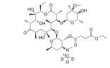


Size: 10 mM × 1 mL, 200 mg

Erythromycin ethylsuccinate-13C,d3

(Erythromycin ethyl succinate-13C,d3; EES-13C,d3)

Erythromycin ethylsuccinate-13C,d3 is the 13C- and deuterium labeled. Erythromycin Ethylsuccinate is an antibiotic useful for the treatment of a number of bacterial infections, has an antimicrobial spectrum similar to or slightly wider than that of penicillin.



Cat. No.: HY-B0957S

Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg

Esmolol hydrochloride

Cat. No.: HY-B1392

Esmolol hydrochloride is a beta adrenergic receptor blocker.

Purity: 99.34% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg

Esmolol-d7 hydrochloride

Esmolol-d7 hydrochloride is the deuterium labeled Esmolol hydrochloride. Esmolol hydrochloride is a beta adrenergic receptor blocker.



Cat. No.: HY-B1392S

Purity: >98% Clinical Data:

Size: 1 mg, 10 mg

Ethyl 3,4-dihydroxybenzoate

(Ethyl protocatechuate)

Cat. No.: HY-W016409

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 Reported Size: 10 mM × 1 mL, 500 mg

Etoposide

(VP-16; VP-16-213)

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**.



Cat. No.: HY-13629

Purity: 99.94% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 200 mg, 500 mg

Etoposide phosphate

(BMY-40481)

Cat. No.: HY-13630

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: 98.40% Clinical Data: Launched

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Etoposide phosphate disodium

(BMY-40481 disodium)

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.



Cat. No.: HY-13630A

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Etoposide-13C,d3

(VP-16-13C,d3; VP-16-213-13C,d3)

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.



Cat. No.: HY-13629S1

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Eupatilin

Eupatilin, a lipophilic flavonoid isolated from Artemisia species, is a **PPAR** α agonist, and possesses anti-apoptotic, anti-oxidative and anti-inflammatory activities.



Cat. No.: HY-N0783

Purity: 98.49% Clinical Data: Phase 4

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg

Everolimus

(RAD001; SDZ-RAD)

Cat. No.: HY-10218

Everolimus (RAD001) is a Rapamycin derivative and a potent, selective and orally active mTOR1 inhibitor. Everolimus binds to FKBP-12 to generate an immunosuppressive complex. Everolimus inhibits tumor cells proliferation and induces cell apoptosis and autophagy.



Purity: 99.74%
Clinical Data: Launched

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Everolimus-d4

(RAD001-d4; SDZ-RAD-d4)

Everolimus-d4 (RAD001-d4) is the deuterium labeled Everolimus. Everolimus (RAD001) is a Rapamycin derivative and a potent, selective and orally active mTOR1 inhibitor. Everolimus binds to FKBP-12 to generate an immunosuppressive complex.



Cat. No.: HY-10218S

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 10 mg

Ezetimibe

(SCH 58235) Cat. No.: HY-17376

Ezetimibe (SCH 58235) is a potent cholesterol absorption inhibitor. Ezetimibe is a Niemann-Pick C1-like1 (NPC1L1) inhibitor, and is a potent Nrf2 activator.

HOP

Purity: 99.93% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg, 500 mg

Ezetimibe ketone

(EZM-K) Cat. No.: HY-133114

Ezetimibe ketone (EZM-K) is a phase-I metabolite of Ezetimibe. Ezetimibe is a Niemann-Pick C1-like1 (NPC1L1) inhibitor, and is a potent Nrf2 activator. Ezetimibe is a potent cholesterol absorption inhibitor.



Purity: >98%

Clinical Data: No Development Reported

Size: 5 mg

Ezetimibe-d4 diacetate

Cat. No.: HY-17376S2

Ezetimibe-d4 diacetate is the deuterium labeled Ezetimibe. Ezetimibe (SCH 58235) is a potent cholesterol absorption inhibitor. Ezetimibe is a Niemann-Pick C1-like1 (NPC1L1) inhibitor, and is a potent Nrf2 activator.

Purity: > 98%

Clinical Data: No Development Reported

Size: 1 mg, 10 mg

Ezetimibe-d4-1

(SCH 58235-d4-1) Cat. No.: HY-17376S1

Ezetimibe-d4 is deuterium labeled Ezetimibe. Ezetimibe (SCH 58235) is a potent cholesterol absorption inhibitor. Ezetimibe is a Niemann-Pick C1-like1 (NPC1L1) inhibitor, and is a potent Nrf2 activator.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

FAAH inhibitor 1

(Benzothiazole analog 3) Cat. No.: HY-10862

FAAH inhibitor 1 (Benzothiazole analog 3) is a potent fatty acid amide hydrolase (FAAH) inhibitor with an IC_{sn} of 18 ± 8 nM.

D1-01-01-6

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

FAAH-IN-1

Cat. No.: HY-111389

FAAH-IN-1 is a fatty acid amide hydrolase (FAAH) inhibitor, with $\rm IC_{50}S$ of 145 nM and 650 nM for rat and human FAAH, respectively.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

FAAH-IN-2

(O-Desmorpholinopropyl Gefitinib) Cat. No.: HY-79511

FAAH-IN-2 (O-Desmorpholinopropyl Gefitinib) is a potent FAAH(fatty acid amide hydrolase) inhibitor extracted from Patent WO/2008/100977A2.



Purity: 98.17%

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g

FAK-IN-5

Cat. No.: HY-147520

FAK-IN-5 (Compound 8I) is a **FAK signaling** inhibitor. FAK-IN-5 induces cell **apoptosis** and **autophagy**.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Falcarindiol

Cat. No.: HY-N0364

Falcarindiol, an orally active polyacetylenic oxylipin, activates PPAR γ and increases the expression of the cholesterol transporter ABCA1 in cells. Falcarindiol induces **apoptosis** and **autophagy**.



Purity: > 98%

Clinical Data: No Development Reported

Size: 5 mg, 10 mg

Fangchinoline

Cat. No.: HY-N1372A

Fangchinoline is isolated from Stephania tetrandra with extensive biological activities, such as enhancing immunity, anti-inflammatory sterilization and anti-atherosclerosis.



Purity: 99.92%

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 5 mg

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Fascaplysin

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).

Cat. No.: HY-112328

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Fasudil Hydrochloride

(HA-1077 Hydrochloride; AT-877 Hydrochloride)

AT877 Hydrochloride), is a nonspecific RhoA/ROCK inhibitor and also has inhibitory effect on protein kinases, with an K_i of 0.33 μM for ROCK1, IC_{so} s of 0.158 μM and 4.58 μM, 12.30 μM, 1.650 μM for ROCK2 and PKA, PKC, PKG, respectively.

Purity: Clinical Data: Launched

10 mM × 1 mL, 200 mg, 500 mg Size:

Fasudil

(HA-1077; AT877)

Fasudil (HA-1077; AT877), is a nonspecific RhoA/ROCK inhibitor and also has inhibitory effect on protein kinases, with an K, of 0.33 µM for ROCK1, IC_{so} s of 0.158 μ M and 4.58 μ M, 12.30 $\mu\text{M},\,1.650~\mu\text{M}$ for ROCK2 and PKA, PKC, PKG, respectively.

Purity: >98% Clinical Data: Launched Size: 1 mg, 5 mg

Cat. No.: HY-10341A

Cat. No.: HY-10341

Fasudil Hydrochloride (HA-1077 Hydrochloride;

99 91%

Felodipine

Felodipine, a dihydropyridine, is a potent, vasoselective calcium channel antagonist. Felodipine lowers blood pressure (BP) by selective action on vascular smooth muscle, especially in the resistance vessels.

Purity: 98 93% Clinical Data: Launched

Felodipine-d5

10 mM × 1 mL, 10 mg, 50 mg



Cat. No.: HY-B0309

Felodipine-d3

Cat. No.: HY-B0309S2

Felodipine-d3 is the deuterium labeled Felodipine. Felodipine, a dihydropyridine, is a potent, vasoselective calcium channel antagonist. Felodipine lowers blood pressure (BP) by selective action on vascular smooth muscle, especially in the resistance vessels.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

H-CI

Felodipine-d5 is deuterium labeled Felodipine. Felodipine, a dihydropyridine, is a potent, vasoselective calcium channel antagonist. Felodipine lowers blood pressure (BP) by selective action on vascular smooth muscle, especially in the resistance vessels.

Purity: Clinical Data:

Size 1 mg, 5 mg



Cat. No.: HY-B0309S1

Felodipine-d8

Cat. No.: HY-B0309S Felodipine-d8 is the deuterium labeled Felodipine.

Felodipine, a dihydropyridine, is a potent, vasoselective calcium channel antagonist. Felodipine lowers blood pressure (BP) by selective action on vascular smooth muscle, especially in the resistance vessels.

Purity: >98%

Clinical Data: No Development Reported

Size: 10 ma



Fenofibrate

Fenofibrate is a selective $PPAR\alpha$ agonist with an EC_{so} of 30 μM. Fenofibrate also inhibits human cytochrome P450 isoforms, with **IC**_{so}s of 0.2, 0.7, 9.7, 4.8 and 142.1 μM for CYP2C19, CYP2B6, CYP2C9, CYP2C8, and CYP3A4, respectively.

Cat. No.: HY-17356

99.92% Purity: Clinical Data: Launched

10 mM \times 1 mL, 200 mg, 5 g, 10 g Size:

Fenofibrate-d6

Cat. No.: HY-17356S

Fenofibrate-d6 is the deuterium labeled Fenofibrate. Fenofibrate is a selective PPARα agonist with an EC_{so} of 30 μM.

Purity: >98%

Clinical Data: No Development Reported

Size 1 mg, 5 mg

Fenretinide

(4-HPR)

Fenretinide (4-HPR) is a synthetic retinoid deriverative, binding to the retinoic acid receptors (RAR) at concentrations necessary to induce cell death.



Cat. No.: HY-15373

Purity: 99.08% Clinical Data: Launched

 $10 \text{ mM} \times 1 \text{ mL}$, 10 mg, 50 mg, 100 mg

Ferulic acid methyl ester

(Methyl ferulate) Cat. No.: HY-W018643

Ferulic acid methyl ester (Methyl ferulate) is a derivative of ferulic acid, isolated from Stemona tuberosa, with anti-inflammatory and antioxidant properties.

Purity: 99.95%

Clinical Data: No Development Reported Size: 10 mM × 1 mL, 100 mg

Fexaramine

Fexaramine is a potent and selective FXR agonist with an EC $_{so}$ of 25 nM. Fexaramine has no activity against hRXR α , hPPAR $\alpha\gamma\delta$, mPXR, hPXR, hLXR α , hTR β , hRAR β , mCAR, mERR γ , and hVDR receptors.



Cat. No.: HY-10912

Purity: 99.29%

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 10 mg, 50 mg

Flavopiridol

(HMR-1275; Alvocidib; L86-8275)

Flavopiridol (Alvocidib) is a broad spectrum and competitive inhibitor of CDKs, inhibiting CDK1, CDK2, CDK4 with $\rm IC_{s0}s$ of 30, 170, 100 nM, respectively.



Cat. No.: HY-10005

Purity: 99.72% Clinical Data: Phase 2

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

FIPI

(5-Fluoro-2-indolyl deschlorohalopemide)

FIPI is a derivative of halopemide which potently inhibits both PLD1 and PLD2 with IC_{50} s of 25 nM and 20 nM, respectively.

HN HN H

Cat. No.: HY-12807

Purity: 99.49%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Flavopiridol Hydrochloride (Alvocidib Hydrochloride; L86-8275

Hydrochloride; HMR-1275 Hydrochloride) Cat. No.: HY-10006

Flavopiridol Hydrochloride (Alvocidib Hydrochloride) is a broad inhibitor of CDK, competing with ATP to inhibit CDKs including CDK1, CDK2, CDK4 with $\rm IC_{50}s$ of 30, 170, 100 nM, respectively.



Purity: 98.95% Clinical Data: Phase 2

Size: $10 \text{ mM} \times 1 \text{ mL}$, 5 mg, 10 mg, 50 mg, 100 mg

Fludrocortisone acetate

(9α-Fludrocortisone acetate; 9α-Fluorcortisol acetate) Cat. No.: HY-B1203A

Fludrocortisone acetate (9α -Fludrocortisone acetate) is a synthetic mineralocorticoid, used to control the amount of sodium and fluids in your body.



Purity: 99.45% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg

Fludrocortisone acetate-d5 (9α-Fludrocortisone acetate-d5;

9α-Fluorcortisol acetate-d5) Cat. No.: HY-B1203AS

Fludrocortisone acetate-d5 is deuterium labeled Fludrocortisone acetate.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Fluvastatin

(XU 62-320 free acid)

Fluvastatin (XU 62-320 free acid) is a first fully synthetic, competitive HMG-CoA reductase inhibitor with an IC $_{\rm 50}$ of 8 nM. Fluvastatin protects vascular smooth muscle cells against oxidative stress through the Nrf2-dependent antioxidant pathway.

Purity: >98%
Clinical Data: Launched
Size: 1 mg, 5 mg



Cat. No.: HY-14664

Fluvastatin D6 sodium

(XU 62-320 (D6)) Cat. No.: HY-14664AS

Fluvastatin D6 sodium (XU 62-320 D6) is deuterium labeled Fluvastatin sodium. Fluvastatin sodium (XU 62320) is a first fully synthetic, competitive $\rm HMG-CoA\ reductase\ inhibitor\ with\ an\ IC_{50}$ of 8 nM.



Purity: > 98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Fluvastatin sodium

(XU 62-320) Cat. No.: HY-14664A

Fluvastatin sodium (XU 62320) is a first fully synthetic, competitive HMG-CoA reductase inhibitor with an IC $_{50}$ of 8 nM. Fluvastatin sodium protects vascular smooth muscle cells against oxidative stress through the Nrf2-dependent antioxidant pathway.

n muscle cells against the Nrf2-dependent

Purity: 99.90% Clinical Data: Launched

Size: 10 mM × 1 mL, 50 mg, 100 mg

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FMK 9a

Cat. No.: HY-100522

FMK 9a is an autophagin-1 inhibitor with IC_{so} values of 80 and 73 μM in FRET and LRA assay.



>98.0% Purity:

FR 167653

(FR 167653 sulfate)

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

FR 167653 (FR 167653 sulfate), an orally active and selective p38 MAPK inhibitor, is a potent suppressor of TNF- α and IL-1 β production via specific inhibition of p38 MAPK activity.

Cat. No.: HY-18754A

Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg

Fulvestrant

(ICI 182780; ZD 9238; ZM 182780) Cat. No.: HY-13636

Fulvestrant (ICI 182780) is a pure antiestrogen and a potent estrogen receptor (ER) antagonist with an IC_{so} of 9.4 nM. Fulvestrant is also a GPR30 agonist. Fulvestrant effectively inhibits the growth of ER-positive MCF-7 cells with an IC_{EQ} of 0.29 nM.



Purity: 99 95% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg

Galangin

(Norizalpinin; 3,5,7-Trihydroxyflavone) Cat. No.: HY-N0382

Galangin (Norizalpinin) is an agonist/antagonist of the arylhydrocarbon receptor. Galangin (Norizalpinin) also shows inhibition of CYP1A1 activity.



Purity: 99.96%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 10 mg, 25 mg, 50 mg, 100 mg

Ganoderic acid A

Cat. No.: HY-N1447

Ganoderic acid A can inhibit of the JAK-STAT3 signaling pathway, also inhibit proliferation, viability, ROS.



Purity: 99.84%

No Development Reported Clinical Data:

Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 25 mg

Forskolin

(Coleonol; Colforsin)

Forskolin (Coleonol) is a potent adenylate cyclase activator with an IC_{50} of 41 nM and an EC_{50} of 0.5 µM for type I adenylyl cyclase. Forskolin is also an inducer of intracellular cAMP

99.82% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg



Cat. No.: HY-15371

FR 167653 free base

FR 167653 free base, an orally active and selective p38 MAPK inhibitor, is a potent suppressor of TNF- α and IL-1 β production via specific inhibition of p38 MAPK activity.

Cat. No.: HY-18754

Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg

Fulvestrant-d3

(ICI 182780-d3; ZD 9238-d3; ZM 182780-d3) Cat. No.: HY-13636S

Fulvestrant-d3 (ICI 182780-d3) is the deuterium labeled Fulvestrant. Fulvestrant (ICI 182780) is a pure antiestrogen and a potent estrogen receptor (ER) antagonist with an IC₅₀ of 9.4 nM. Fulvestrant is also a GPR30 agonist.



>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Gambogic Acid

(Beta-Guttiferrin)

Gambogic Acid (Beta-Guttiferrin) is derived from the gamboges resin of the tree Garcinia hanburyi. Gambogic Acid (Beta-Guttiferrin) inhibits Bcl-X,, Bcl-2, Bcl-W, Bcl-B, Bfl-1 and Mcl-1 with ICsos of 1.47 $\mu\text{M},\,1.21~\mu\text{M},\,2.02~\mu\text{M},\,0.66~\mu\text{M},\,1.06~\mu\text{M}$ and $0.79 \mu M.$



Cat. No.: HY-N0087

Purity: 95.27%

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

GANT 61

(NSC 136476)

GANT 61 is an inhibitor of Gli1 and Gli2 targeting the Hedgehog/GLI pathway.



Cat. No.: HY-13901

≥98.0% **Purity:**

Clinical Data: No Development Reported 10 mM × 1 mL, 5 mg, 10 mg, 50 mg

Gartanin

Cat. No.: HY-N6038

Gartanin is a natural xanthone of mangosteen, with antioxidant, anti-inflammatory, antifungal, neuroprotective and antineoplastic properties. Gartanin induces cell cycle arrest and autophagy and suppresses migration in human glioma cells.



Purity: >97.0%

Clinical Data: No Development Reported

Size: 5 mg

GDC-0349

Cat. No.: HY-15248

GDC-0349 is a potent and selective ATP-competitive mTOR inhibitor with a K_i of 3.8 nM. GDC-0349 inhibits of both mTORC1 and mTORC2 complexes.



Purity: 98 42% Clinical Data: Phase 1

10 mM × 1 mL, 5 mg, 10 mg, 50 mg

GCN2iB

GCN2iB is an ATP-competitive inhibitor of a serine/threonine-protein kinase general control nonderepressible 2 (GCN2), with an IC₅₀ of 2.4



Cat. No.: HY-112654

99.81% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Gefitinib

(ZD1839) Cat. No.: HY-50895

Gefitinib (ZD1839) is a potent, selective and orally active EGFR tyrosine kinase inhibitor with an IC_{so} of 33 nM. Gefitinib selectively inhibits EGF-stimulated tumor cell growth (IC₅₀ of 54 nM) and that blocks EGF-stimulated EGFR autophosphorylation in tumor cells.



99.94% **Purity:** Clinical Data: Launched

10 mM × 1 mL, 100 mg, 500 mg, 1 g, 5 g Size:

Gefitinib-d3

Cat. No.: HY-50895S2

Gefitinib-d3 (ZD1839-d3) is the deuterium labeled Gefitinib. Gefitinib (ZD1839) is a potent, selective and orally active EGFR tyrosine kinase inhibitor with an IC₅₀ of 33 nM.



>98% Purity:

Clinical Data:

Size: 1 mg, 10 mg

Gefitinib-d6

(ZD1839-d6) Cat. No.: HY-50895S1

Gefitinib-d6 (ZD1839-d6) is the deuterium labeled Gefitinib. Gefitinib (ZD1839) is a potent, selective and orally active EGFR tyrosine kinase inhibitor with an IC_{50} of 33 nM.



>98% Purity:

Clinical Data: No Development Reported

Size 1 mg, 5 mg

Gefitinib-d8

(ZD1839-d8) Cat. No.: HY-50895S

Gefitinib D8 (ZD1839 D8) is a deuterium labeled Gefitinib. Gefitinib is an EGFR tyrosine kinase inhibitor, with IC_{so} of 2-37 nM in NR6wtEGFR cells.



98.42% Purity:

Clinical Data: No Development Reported

Size 5 ma

Gemcitabine

(LY 188011) Cat. No.: HY-17026

Gemcitabine (LY 188011) is a pyrimidine nucleoside analog antimetabolite and an antineoplastic agent. Gemcitabine inhibits DNA synthesis and repair, resulting in autophagyand apoptosis.



99.92% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 200 mg, 500 mg, 1 g

Gemcitabine elaidate

(CP-4126; CO-101; Gemcitabine 5'-elaidate) Cat. No.: HY-13538

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: 98.22% Clinical Data: Launched

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Gemcitabine elaidate hydrochloride (CP-4126 hydrochloride;

CO-101 hydrochloride; ...) Cat. No.: HY-13538A

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.



Purity: ≥97.0%

Clinical Data: No Development Reported 5 mg, 10 mg, 50 mg, 100 mg

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Gemcitabine hydrochloride

(LY 188011 hydrochloride) Cat. No.: HY-B0003

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: 99.93% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 200 mg, 500 mg, 1 g

Genipin

((+)-Genipin) Cat. No.: HY-17389

Genipin ((+)-Genipin) is a natural crosslinking reagent derived from Gardenia jasminoides Ellis fruits. Genipin inhibits UCP2 (uncoupling protein 2) in cells.

Purity: 99.40%

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



Genistein

(NPI 031L) Cat. No.: HY-14596

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-d4

(NPI 031L-d4) Cat. No.: HY-14596S

Genistein-d4 (NPI 031L-d4) is the deuterium labeled Genistein. Genistein, a soy isoflavone, is a multiple **tyrosine kinases** (e.g.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Ginkgetin

Cat. No.: HY-N0889

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_{τ_0} of 5.92 μ M.

Purity: 99.53%

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 5 mg, 10 mg

Ginkgolide K

Ginkgolide K, isolated from Ginkgo biloba, induces protective autophagy through the AMPK/mTOR/ULK1 signaling pathway. Ginkgolide K possesses neuroprotective activity.



Cat. No.: HY-N4176

Purity: 99.85%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 25 mg

Ginsenoside F2

Cat. No.: HY-125848

Ginsenoside F2, a metabolite from Ginsenoside Rb1, induces apoptosis accompanied by protective autophagy in breast cancer stem cells.

Purity: 99.95%

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 5 mg, 10 mg, 20 mg

Ginsenoside Rb1

(Gypenoside III)

Ginsenoside Rb1, a main constituent of the root of Panax ginseng, inhibits $Na^{\star}, K^{\star}\text{-ATPase}$ activity with an IC_{so} of $6.3\pm1.0~\mu\text{M}.$ Ginsenoside also inhibits IRAK-1 activation and phosphorylation of NF- κB p65 .



Cat. No.: HY-N0039

Purity: 98.75%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Glaucocalyxin B

Cat. No.: HY-N2113

Glaucocalyxin B is an ent kaurane diterpenoid isolated from the Chinese traditional medicine Rabdosia japonica with anticancer and antitumor activity; decreases the growth of HL-60 cells with an ${\rm IC}_{sn}$ of approximately 5.86 μ M at 24 h.

Purity: 99.39%

Clinical Data: No Development Reported Size: 10 mM × 1 mL, 1 mg, 5 mg

Glibenclamide

(Glyburide)

Glibenclamide (Glyburide) is an orally active ATP-sensitive K* channel ($K_{\rm ATP}$) inhibitor and can be used for the research of diabetes and obesity. Glibenclamide inhibits **P-glycoprotein**.



Cat. No.: HY-15206

Purity: 99.79% Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g

GLPG1837

(ABBV-974) Cat. No.: HY-111099

GLPG1837 is a potent and reversible CFTR potentiator, with EC_{so}s of 3 nM and 339 nM for F508del and G551D CFTR, respectively.

Purity: 99.03%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 50 mg, 100 mg

Glucosamine

(D-Glucosamine; Chitosamine)

Glucosamine (D-Glucosamine) is an amino sugar and a prominent precursor in the biochemical synthesis of glycosylated proteins and lipids, is used as a dietary supplement.

Cat. No.: HY-B1125

Purity: > 98.0% Clinical Data: Launched 100 ma Size:

Glucosamine sulfate

(D-Glucosamine sulfate)

Glucosamine sulfate (D-Glucosamine sulfate) is an amino sugar and a prominent precursor in the biochemical synthesis of glycosylated proteins and lipids, is used as a dietary supplement.

Cat. No.: HY-N0487

Purity: ≥98.0% Clinical Data: Launched Size: 500 ma

Glyburide-d11

Cat. No.: HY-15206S Glyburide-d11 is the deuterium labeled

Glibenclamide. Glibenclamide (Glyburide) is an orally active ATP-sensitive K+ channel (K_{ATP}) inhibitor and can be used for the research of diabetes and obesity. Glibenclamide inhibits P-glycoprotein.

Purity: >98%

Clinical Data: No Development Reported

1 mg, 10 mg Size:

Cat. No.: HY-N8034

Glychionide A

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.

Purity: >98%

Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg

GLPG2451

GLPG2451 is a cystic fibrosis transmembrane conductance regulator (CFTR) potentiator, which effectively potentiates low temperature rescued F508del CFTR with an EC₅₀ of 11.1 nM.

Cat. No.: HY-119936

Purity: 99 62%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Glucosamine hydrochloride (D-(+)-Glucosamine hydrochloride;

Chitosamine hydrochloride)

Glucosamine hydrochloride (D-Glucosamine hydrochloride) is an amino sugar and a prominent precursor in the biochemical synthesis of glycosylated proteins and lipids, is used as a dietary supplement.

Purity: ≥98.0% Clinical Data: Launched

10 mM × 1 mL, 500 mg

Cat. No.: HY-N0733

Gly-β-MCA

Gly-β-MCA, a bile acid, is a potent, sable, intestine-selective and oral bioactive farnesoid X receptor (FXR) inhibitor that may be a candidate for the treatment of metabolic disorders.

Cat. No.: HY-15206S1

Cat. No.: HY-114392

98.11% Purity:

Clinical Data: No Development Reported Size 10 mM × 1 mL, 5 mg, 10 mg

Glyburide-d3

(Glyburide-d3)

Glyburide-d3 (Glyburide-d3) is the deuterium labeled Glibenclamide. Glibenclamide (Glyburide) is an orally active ATP-sensitive K+ channel (K_{ATP}) inhibitor and can be used for the research of diabetes and obesity. Glibenclamide inhibits P-glycoprotein.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Glycycoumarin

Glycycoumarin is a major bioactive coumarin of licorice. Glycycoumarin inhibits hepatocyte lipoapoptosis through activation of autophagy and inhibition of ER stress-mediated JNK and GSK-3-mediated mitochondrial pathway.



Cat. No.: HY-N4113

Purity: 99.89%

Clinical Data: No Development Reported 10 mM × 1 mL, 5 mg, 10 mg

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Glyphosate

Cat. No.: HY-B0863

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.

Purity: >98.0%

Clinical Data: No Development Reported Size: $10 \text{ mM} \times 1 \text{ mL}, 500 \text{ mg}$

Glyphosate-d2

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.



Cat. No.: HY-10249

Cat. No.: HY-B0863S

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

GPP78

(CAY10618) Cat. No.: HY-14374

GPP78 (CAY10618) is a potent Nampt inhibitor with an IC_{50} of 3.0 nM for nicotinamide adenine dinucleotide (NAD) depletion. GPP78 is cytotoxic to neuroblastoma cell line SH-SY5Y cells with an IC₅₀ of 3.8 nM by inducing autophagy. GPP78 has anti-cancer and anti-inflammatory effects.



Purity:

Clinical Data: No Development Reported

GSK-690693

GSK-690693 is an ATP-competitive pan-Akt inhibitor with IC₅₀s of 2 nM, 13 nM, 9 nM for Akt1, Akt2 and Akt3, respectively. GSK-690693 is also an AMPK inhibitor, affects Unc-51-like autophagy activating kinase 1 (ULK1) activity and robustly inhibits STING-dependent IRF3 activation.

Purity: 98 40% Clinical Data: Phase 1

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

5 mg (11.38 mM * 1 mL in Methanol),

GSK2578215A

Cat. No.: HY-13237

GSK2578215A is a potent and highly selective LRRK2 inhibitor, which exhibits IC₅₀s of around 10 nM against both wild-type LRRK2 and the G2019S mutant.



Purity: 99.79%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

GSK2606414

Cat. No.: HY-18072

GSK2606414 is a cell-permeable and orally available protein kinase R-like endoplasmic reticulum (ER) kinase (PERK) inhibitor with an IC₅₀ of 0.4 nM.



99.89% Purity:

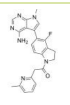
Clinical Data: No Development Reported

Size 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

GSK2656157

Cat. No.: HY-13820

GSK2656157 is a selective and ATP-competitive inhibitor of protein kinase R (PKR)-like endoplasmic reticulum kinase (PERK) with an ${\rm IC}_{\rm co}$ of 0.9 nM.



99.66% Purity:

Clinical Data: No Development Reported

10 mM \times 1 mL, 10 mg, 50 mg, 100 mg Size:

GSK343

Cat. No.: HY-13500

GSK343 is a highly potent and selective EZH2 inhibitor with an IC_{50} of 4 nM.



99.45% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

GSK4112

(SR6452) Cat. No.: HY-14414

GSK4112 is a Rev-erbα agonist with EC50 of 0.4 μM, also is a small molecule chemical probe for the cell biology of the nuclear heme receptor Rev-erbα.



Purity: ≥98.0%

Clinical Data: No Development Reported

Size 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

GSK621

Cat. No.: HY-100548

GSK621 is a specific AMPK activator, with IC_{so} values of 13-30 μM for AML cells. GSK621 induces autophagy and apoptosis. GSK621 induces eiF2α phosphorylation-a hallmark of UPR activation.



Purity: 98.82%

Clinical Data: No Development Reported 5 mg, 10 mg, 25 mg

Guaiol

(Champacol; Guaiac alcohol)

Guaiol is a sesquiterpene alcohol that has been found in several traditional Chinese medicinal plants and has antiproliferative, pro-autophagic, insect repellent, and insecticidal biological activities.



98 67%

Clinical Data: No Development Reported

Size: 5 mg, 10 mg

Cat. No.: HY-N3980

Purity:

Purity:

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Guggulsterone is a plant sterol derived from the

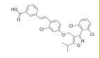
gum resin of the tree Commiphora wightii.

99.83% Clinical Data: No Development Reported

GW 4064

Cat. No.: HY-50108

GW 4064 is a potent FXR agonist with an EC_{so} of 65 nM.



Purity: 99 76%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

GW 501516

Guggulsterone

(Z/E-Guggulsterone)

(GW 1516; GSK-516) Cat. No.: HY-10838

GW 501516 (GW 1516) is a PPARδ agonist with an EC₅₀ of 1.1 nM.



Cat. No.: HY-N2541

Cat. No.: HY-107738

Purity: 99 15% Clinical Data: Phase 4

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

GW406108X

(GW108X) Cat. No.: HY-115570

GW406108X is a specific Kif15 (Kinesin-12) inhibitor with an IC₅₀ of 0.82 uM in ATPase assays. GW406108X, a potent autophagy inhibitor, shows ATP competitive inhibition against ULK1 with a pIC₅₀ of 6.37 (427 nM).

96.02% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Gymnemic acid I

Gymnemic acid I is a bioactive triterpene saponin found in Gymnema sylvestre. Gymnemic acid I decreases the apoptosis under the high glucose

Purity:

96.31%

Clinical Data: No Development Reported

Size: 1 ma, 5 ma

H-89

Cat. No.: HY-15979

H-89 is a potent and selective inhibitor of cyclic AMP-dependent protein kinase (protein kinase A) with IC₅₀ of 48 nM and has weak inhibition on PKG, PKC, Casein Kinase, and others kinases.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

H-89 dihydrochloride

Cat. No.: HY-15979A

H-89 dihydrochloride is a potent and selective inhibitor of protein kinase A (PKA) with an IC₅₀ of 48 nM and has weak inhibition on PKG, PKC, Casein Kinase.



99.34% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg

HA15

Cat. No.: HY-100437

HA15 is a potent and specific inhibitor of ER chaperone BiP/GRP78/HSPA5, inhibits the ATPase activity of BiP, with anti-cancerous activity.



Purity: 99.62%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

HDAC-IN-36

HDAC-IN-36 (compound 23 g) is an orally active and

potent HDAC (histone deacetylase) inhibitor, with an IC_{so} of 11.68 nM (HDAC6). HDAC-IN-36 promotes apoptosis, autophagy and suppresses migration.



Cat. No.: HY-146684

>98% Purity:

Clinical Data: No Development Reported

1 mg, 5 mg

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

Hemin

(Hemin chloride) Cat. No.: HY-19424

Hemin is an iron-containing porphyrin. Hemin is an Heme oxygenase (HO)-1 inducer.



>98% Purity: Clinical Data: Launched

Size: $10 \text{ mM} \times 1 \text{ mL}, 500 \text{ mg}, 1 \text{ g}, 5 \text{ g}$

Heparin

Heparin is a highly sulfated

glycosaminoglycan, that is widely used as an injectable anticoagulant, and has the highest negative charge density of any known biological molecule. Heparin significantly inhibits exosome-cell interactions.

Purity: >98% Clinical Data: Launched

10 mg(10 mg × mL in Water) Size:



Cat. No.: HY-17567

Heparin Lithium salt

Cat. No.: HY-17567B

Heparin Lithium salt is an anticoagulant which binds reversibly to antithrombin III (ATIII). Heparin Lithium salt significantly inhibits exosome-cell interactions.



Purity: >98% Clinical Data: Launched

10 mg(10 mg × mL in Water), 100 mg, 500 mg Size:

Heparin sodium salt

(Sodium heparin; Sodium heparinate)

Heparin sodium salt (Sodium heparin) is an anticoagulant which binds reversibly to antithrombin III (ATIII) and greatly accelerates the rate at which ATIII inactivates coagulation enzymes thrombin factor IIa and factor Xa.



Cat. No.: HY-17567A

Purity: >98% Clinical Data: Launched 100 mg, 500 mg, 1 g

Heparin sodium salt (MW 15kDa)

(Sodium heparin (MW 15kDa); Sodium heparinate (MW 15kDa) ht. No.: HY-17567C

Heparin sodium salt (MW 15kDa) (Sodium heparin (MW 15kDa)) is a polymer of Heparin with the molecular weight of 15kDa.



Purity: >98% Clinical Data: Launched Size: 100 mg, 500 mg

Hesperadin

Cat. No.: HY-12054

Hesperadin is an ATP competitive indolinone inhibitor of Aurora A and B. Hesperadin inhibits Aurora B with an IC₅₀ of 250 nM. Hesperadin inhibits the growth of Trypanosoma brucei by blocking nuclear division and cytokinesis.

Purity: ≥98.0%

Clinical Data: No Development Reported

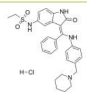
Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg



Hesperadin hydrochloride

Cat. No.: HY-12054A

Hesperadin hydrochloride is an ATP competitive indolinone inhibitor of Aurora A and B. Hesperadin hydrochloride inhibits Aurora B with an IC₅₀ of 250 nM.



>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Hesperetin

Hesperetin is a natural flavanone, and acts as a potent and broad-spectrum inhibitor against human

UGT activity. Hesperetin induces apoptosis.



Cat. No.: HY-N0168

98.75% Purity:

Clinical Data: No Development Reported 10 mM × 1 mL, 50 mg Size:

Hesperidin

(Hesperetin 7-rutinoside) Cat. No.: HY-15337

Hesperidin (Hesperetin 7-rutinoside), a flavanone glycoside, is isolated from citrus fruits. Hesperidin has numerous biological properties, such as decreasing inflammatory mediators and exerting significant antioxidant effects.



Purity: 99.19% Clinical Data: Launched

Size 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg, 500 mg, 1 g

HMG499

HMG499 is a potent and selective HMG-CoA reductase inhibitor with an IC_{so} of 0.41 μM .

HMG499 can prevent statins-induced accumulation of HMGCR, reduce serum cholesterol levels and

decrease atherosclerosis.

Purity: >98%

Clinical Data: No Development Reported

5 mg, 10 mg



Cat. No.: HY-114316

Hoechst 33342

(bisBenzimide H 33342; HOE 33342)

Cat. No.: HY-15559

Hoechst 33342 is a DNA minor groove binder used fluorochrome for visualizing cellular **DNA**.

Purity: 99.24%

Clinical Data: No Development Reported

Size: 50 mg, 100 mg

Hoechst 33342 trihydrochloride (bisBenzimide H 33342

trihydrochloride; HOE 33342 trihydrochloride)

Hoechst 33342 trihydrochloride is a membrane permeant blue fluorescent **DNA** stain.

Cat. No.: HY-W031727

Cat. No.: HY-15559A

Purity: 99.87%

Clinical Data: No Development Reported

Size: 50 mg, 100 mg

Honokiol

(NSC 293100) Cat. No.: HY-N0003

Honokiol is a bioactive, biphenolic phytochemical that possesses potent antioxidative, anti-inflammatory, antiangiogenic, and anticancer activities by targeting a variety of signaling molecules. It inhibits the activation of Akt.

Purity: 99.90% Clinical Data: Phase 3

Size: 10 mM × 1 mL, 50 mg, 100 mg, 200 mg

Hydroxychloroquine

Hydroxychloroquine is a synthetic **antimalarial** agent which can also inhibit **Toll-like receptor 7/9** (**TLR7/9**) signaling. Hydroxychloroquine is efficiently inhibits **SARS-CoV-2** infection in

vitro.

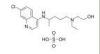
Purity: ≥97.0%
Clinical Data: Launched
Size: 1 mg, 5 mg

>97.0%

Hydroxychloroquine sulfate

(HCQ sulfate) Cat. No.: HY-B1370

Hydroxychloroquine sulfate (HCQ sulfate) is a synthetic antimalarial agent which can also inhibit Toll-like receptor 7/9 (TLR7/9) signaling. Hydroxychloroquine sulfate is efficiently inhibits SARS-CoV-2 infection in vitro.



Purity: 99.99% Clinical Data: Launched

Size: 10 mM × 1 mL, 50 mg

Hydroxychloroquine-d4 sulfate

(HCQ-d4 sulfate) Cat. No.: HY-B1370S

Hydroxychloroquine-d4 sulfate (HCQ-d4 sulfate) is the deuterium labeled Hydroxychloroquine sulfate. Hydroxychloroquine sulfate (HCQ sulfate) is a synthetic antimalarial agent which can also inhibit Toll-like receptor 7/9 (TLR7/9) signaling.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Hydroxychloroquine-d4-1 sulfate

Cat. No.: HY-W031727S

Hydroxychloroquine-d4-1 sulfate is the deuterium labeled Hydroxychloroquine. Hydroxychloroquine is a synthetic antimalarial agent which can also inhibit Toll-like receptor 7/9 (TLR7/9) signaling. Hydroxychloroquine is efficiently inhibits SARS-CoV-2 infection in vitro.



Purity: > 98%

Clinical Data: No Development Reported

Size: 1 mg, 10 mg

Hydroxyurea

(Hydroxycarbamide)

Hydroxyurea is a cell apoptosis inducer that inhibit **DNA** synthesis through inhibition of **ribonucleotide reductase**.



Cat. No.: HY-B0313

Purity: ≥98.0% Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g

Icariin

(Ieariline) Cat. No.: HY-N0014

Icariin is a flavonol glycoside. Icariin inhibits PDE5 and PDE4 activities with IC_{so} s of 432 nM and 73.50 μ M, respectively. Icariin also is a PPAR α activator.



Purity: 99.06% Clinical Data: Phase 3

Size: 10 mM × 1 mL, 100 mg, 200 mg, 500 mg

Icaritin

(Anhydroicaritin)

Icaritin (Anhydroicaritin) is a prenylflavonoid derivative from Epimedium Genusis and potently inhibits proliferation of K562 cells (IC $_{50}$ of 8 μ M) and primary CML cells (IC $_{50}$ of 13.4 μ M for CML-CP and 18 μ M for CML-BC).



Cat. No.: HY-N0678

Purity: 99.27% Clinical Data: Phase 2

Size: 10 mM × 1 mL, 5 mg, 10 mg

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

ICCB-19 hydrochloride

Cat. No.: HY-138779

ICCB-19 hydrochloride is a TRADD (TNFRSF1A associated via death domain) inhibitor. ICCB-19 hydrochloride binds with N-terminal domain of TRADD (TRADD-N), disrupting its binding to both TRADD-C and TRAF2. ICCB-19 hydrochloride is indirect inhibitor of RIPK1 kinase activity.

H-CI

Purity: 99 20%

Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Idelalisib D5

Size:

(CAL-101 D5; GS-1101 D5)

Idarubicin hydrochloride

antileukemic drug. It inhibits the

topoisomerase II interfering with the

99 82%

(4-Demethoxydaunorubicin hydrochloride)

Idarubicin hydrochloride is an anthracycline

replication of DNA and RNA transcription.

Idelalisib D5 is a deuterium labeled Idelalisib. Idelalisib is a highly selective and orally bioavailable p110 δ inhibitor.

10 mM × 1 mL, 5 mg, 10 mg, 50 mg

Purity: >98%

Clinical Data: No Development Reported

Idarubicin hydrochloride inhibits the growth of bacteria and yeasts. Purity:

Cat. No.: HY-17381

Idelalisib

(CAL-101; GS-1101)

Idelalisib (CAL-101; GS-1101) is a highly selective and orally bioavailable $p110\delta$ inhibitor with an IC_{so} of 2.5 nM, showing 40- to 300-fold selectivity for p110δ over other PI3K class I enzymes.

Cat. No.: HY-13026

Purity: 99 78% Clinical Data: Launched

10 mM × 1 mL, 10 mg, 25 mg, 50 mg

Clinical Data: Launched

Cat. No.: HY-13026S

IITZ-01

Cat. No.: HY-112897

IITZ-01 is a potent lysosomotropic autophagy inhibitor with single-agent antitumor activity, with an IC_{50} of 2.62 μM for PI3Ky.

99.05% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Imatinib

(STI571; CGP-57148B)

Imatinib (STI571) is an orally bioavailable tyrosine kinases inhibitor that selectively inhibits BCR/ABL, v-Abl, PDGFR and c-kit kinase activity.



Cat. No.: HY-15463

99 54% Purity: Clinical Data: Launched

Size 10 mM × 1 mL, 200 mg, 500 mg, 1 g, 5 g

Imatinib D4

(STI571 D4; CGP-57148B D4)

Imatinib D4 (STI571 D4) is a deuterium labeled Imatinib (STI571). Imatinib is an orally bioavailable tyrosine kinases inhibitor that selectively inhibits BCR/ABL, v-Abl, PDGFR and c-kit kinase activity.

Cat. No.: HY-15463S1

≥99.0% Purity:

Clinical Data: No Development Reported

Size 1 mg, 5 mg

Imatinib Mesylate

(STI571 Mesylate; CGP-57148B Mesylate)

Imatinib Mesylate (STI571 Mesylate) is a tyrosine kinases inhibitor that inhibits c-Kit, Bcr-Abl, and PDGFR (IC_{so}=100 nM) tyrosine kinases.



Cat. No.: HY-50946

99.91% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 200 mg, 500 mg, 1 g, 5 g

Imatinib-d8

(STI571-d8; CGP-57148B-d8)

Imatinib D8 (STI571 D8) is a deuterium labeled Imatinib (STI571). Imatinib is an orally bioavailable tyrosine kinases inhibitor that selectively inhibits BCR/ABL, v-Abl, PDGFR and c-kit kinase activity.



Cat. No.: HY-15463S

Purity: >98%

Clinical Data: No Development Reported

Size

Imiquimod

(R 837)

Imiquimod (R 837), an immune response modifier, is a selective toll like receptor 7 (TLR7) agonist. Imiguimod exhibits antiviral and antitumor effects in vivo. Imiquimod can be used for the research of external genital, perianal warts, cancer and COVID-19.

Purity: 99.96% Clinical Data: Launched

100 mg, 200 mg, 500 mg



Cat. No.: HY-B0180

Imiquimod hydrochloride

(R 837 hydrochloride) Cat. No.: HY-B0180A

Imiquimod hydrochloride (R 837 hydrochloride), an immune response modifier, is a selective toll like receptor 7 (TLR7) agonist. Imiquimod hydrochloride exhibits antiviral and antitumor effects in vivo

HCI

Purity: 99 80% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 200 mg, 500 mg

Imiquimod maleate

(R 837 maleate)

Imiquimod maleate (R 837 maleate), an immune response modifier, is a selective toll like receptor 7 (TLR7) agonist. Imiquimod maleate exhibits antiviral and antitumor effects in vivo.



Cat. No.: HY-B0180B

Purity: >98% Clinical Data: Launched Size: 1 mg, 5 mg

Imiquimod-d6

(R 837-d6) Cat. No.: HY-B0180S

Imiguimod-d6 (R 837-d6) is the deuterium labeled Imiquimod. Imiquimod (R 837), an immune response modifier, is a selective toll like receptor 7 (TLR7) agonist. Imiquimod exhibits antiviral and antitumor effects in vivo.

Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg

Imiquimod-d9

(R 837-d9) Cat. No.: HY-B0180S1

Imiguimod-d9 is deuterium labeled Imiguimod. Imiquimod (R 837), an immune response modifier, is a selective toll like receptor 7 (TLR7) agonist. Imiquimod exhibits antiviral and antitumor effects in vivo.

Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg



Indomethacin

(Indometacin) Cat. No.: HY-14397

Indomethacin (Indometacin) is a potent, blood-brain permeable and nonselective inhibitor of COX1 and COX2, with IC_{so}s of 18 nM and 26 nM for human COX-1 and COX-2, respectively, in CHO cells. Indomethacin disrupts autophagic flux by disturbing the normal functioning of lysosomes.



99.97% Purity: Clinical Data: Launched

10 mM × 1 mL, 500 mg, 1 g, 5 g Size:

Indomethacin farnesil

(Indometacin farnesil) Cat. No.: HY-111274

Indomethacin farnesil is an orally active prodrug of Indomethacin. Indomethacin (Indometacin) is a potent, blood-brain permeable and nonselective inhibitor of COX1 and COX2, with IC₅₀s of 18 nM and 26 nM for human COX-1 and COX-2, respectively, in CHO cells.



Purity: >98%

Clinical Data: No Development Reported

Size 1 mg, 5 mg

Indomethacin sodium hydrate

(Indometacin sodium hydrate) Cat. No.: HY-14397A

Indomethacin sodium hydrate (Indometacin sodium hydrate) is a potent, blood-brain permeable and nonselective inhibitor of COX1 and COX2, with IC₅₀s of 18 nM and 26 nM for human COX-1 and COX-2, respectively, in CHO cells.



96.84% Purity: Clinical Data: Launched

10 mM × 1 mL, 500 mg, 1 g Size:

Indomethacin-d4

(Indometacin-d4) Cat. No.: HY-14397S

Indomethacin-D4 (Indometacin-D4) is a deuterium labeled Indomethacin. Indomethacin is a potent, blood-brain permeable and nonselective inhibitor of COX1 and COX2, with IC_{so}s of 18 nM and 26 nM for human COX-1 and COX-2, respectively, in CHO cells.



Clinical Data: No Development Reported

Size: 1 mg, 5 mg

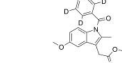


Cat. No.: HY-134807

Indomethacin-d4 Methyl Ester

Cat. No.: HY-14397S1

Indomethacin-d4 Methyl Ester is the deuterium labeled Indomethacin. Indomethacin (Indometacin) is a potent, blood-brain permeable and nonselective inhibitor of COX1 and COX2, with IC_{so}s of 18 nM and 26 nM for human COX-1 and COX-2, respectively, in CHO cells.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 10 mg

Indophagolin

Indophagolin is a potent, indoline-containing autophagy inhibitor (IC₅₀=140 nM). Indophagolin

antagonizes the purinergic receptor P2X, as well as $P2X_1$ and $P2X_3$ with IC_{50} s of 2.71, 2.40 and

3.49 µM, respectively.

Purity: 98.05%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

INT-767

Cat. No.: HY-12434

INT-767 is a dual farnesoid X receptor (FXR)/TGR5 agonist with mean EC_{so}s of 30 and 630 nM, respectively.



>98.0% Purity:

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

Iohexol

Iohexol is a radiographic contrast agent and can be applied for myelography, computerized tomography (cisternography, ventriculography) and MicroCT imaging in vivo.



Cat. No.: HY-18337

Cat. No.: HY-B0594

99 20% Purity: Clinical Data: Phase 4

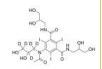
Size:

10 mM × 1 mL, 100 mg, 500 mg

Iohexol-d5

Cat. No.: HY-B0594S

Iohexol-d5 is deuterium labeled Iohexol. Iohexol is a radiographic contrast agent and can be applied for myelography, computerized tomography (cisternography, ventriculography) and MicroCT imaging in vivo.



Purity: >98%

Clinical Data:

Size: 1 mg, 5 mg

IOWH-032

IOWH-032 is a novel and potent CFTR inhibitor (IC50=1.01 uM) in T84 and CHO-CFTR cell based assays. IC50 value: 1.01 uM (CHO-CFTR FLIPR) Target: CFTR Profiling of iOWH032 showed it to be a CFTR inhibitor in T84 and CHO-CFTR cell based

assavs.

Purity: 99.63% Clinical Data: Phase 2

10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 500 mg Size:

Ipsalazide

Cat. No.: HY-101744

Ipsalazide is a novel sulfasalazine analog designed to release 5-aminosalicylic acid and a nontoxic carrier molecule in the gastrointestinal tract.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Irinotecan

((+)-Irinotecan; CPT-11)

Irinotecan ((+)-Irinotecan) is a topoisomerase I inhibitor, preventing religation of the DNA strand by binding to topoisomerase I-DNA complex.



Cat. No.: HY-16562

99.84% Purity: Clinical Data: Launched

Size 10 mM × 1 mL, 50 mg, 100 mg, 200 mg, 500 mg

Irinotecan hydrochloride

((+)-Irinotecan hydrochloride; CPT-11 hydrochloride) Cat. No.: HY-16562A

Irinotecan hydrochloride ((+)-Irinotecan hydrochloride) is a topoisomerase I inhibitor mainly used to treat colon cancer and rectal cancer.



99.75% Purity: Clinical Data: Launched

Size 10 mM × 1 mL, 50 mg, 100 mg, 200 mg, 500 mg

Irinotecan hydrochloride trihydrate ((+)-Irinotecan

hydrochloride trihydrate; ...)

Irinotecan hydrochloride trihydrate ((+)-Irinotecan hydrochloride trihydrate) is a topoisomerase I inhibitor with antitumor

activity.



Cat. No.: HY-16568

99.87% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 50 mg, 100 mg, 200 mg, 500 mg

Irinotecan-d10

((+)-Irinotecan-d10; CPT-11-d10) Cat. No.: HY-16562S

Irinotecan-d10 ((+)-Irinotecan-d10) is a deuterium labeled Irinotecan ((+)-Irinotecan). Irinotecan ((+)-Irinotecan) is a topoisomerase I inhibitor, preventing religation of the DNA strand by binding to topoisomerase I-DNA complex.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Irinotecan-d10 hydrochloride

Cat. No.: HY-16562S1

Irinotecan-d10 ((+)-Irinotecan-d10) hydrochloride is the deuterium labeled Irinotecan. Irinotecan ((+)-Irinotecan) is a topoisomerase I inhibitor, preventing religation of the DNA strand by binding to topoisomerase I-DNA complex.



Purity: >98%

Clinical Data: No Development Reported 1 mg, 5 mg, 10 mg

Isoalantolactone

((+)-Isoalantolactone; Isohelenin) Cat. No.: HY-N0780

Isoalantolactone is an apoptosis inducer, which also acts as an alkylating agent.

Cat. No.: HY-N2585

99 99% Purity:

Isodeoxyelephantopin

isolated from Elephantopus scaber. Isodeoxyelephantopin induces ROS generation,

activities against breast cancer.

Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 50 mg

Isodeoxyelephantopin is a sesquiterpene lactone

suppresses NF-kB activation. Isodeoxyelephantopin

also modulates LncRNA expression and exhibit

Clinical Data: No Development Reported

Isobavachalcone

(Corylifolinin; Isobacachalcone)

Isobavachalcone (Corylifolinin) is derived from Psoralea corvlifolia 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_{50} value of 7.92 μ M).



Cat. No.: HY-13065

99.01% Purity:

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

Cat. No.: HY-19826

Isofistularin-3 is a direct, DNA-competitive DNMT1 inhibitor, with an IC_{s0} of 13.5 $\mu M.$ Isofistularin-3, as a DNA demethylating agent, induces cell cycle arrest and sensitization to TRAIL in cancer cells. Isofistularin-3 can be used

Clinical Data: No Development Reported



Size: 1 mg, 5 mg

(GU17; ISL; Isoliquiritigen)

Purity:

Isoliquiritigenin

Isoliquiritigenin is an anti-tumor flavonoid from the root of Glycyrrhiza glabra, which inhibits aldose reductase with an IC_{so} of 320 nM. Isoliquiritigenin is a potent inhibitor of influenza virus replication with an EC₅₀ of 24.7 μΜ.

Cat. No.: HY-N0102

Purity: 98.17%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg

Isofistularin-3

as an ADC cytotoxin.

Purity: >98%

Size: 1 mg, 5 mg

Isoniazid

(INH; Isonicotinic acid hydrazide; Isonicotinic hydrazide) Cat. No.: HY-B0329

Isoniazid (INH) is a prodrug and must be activated by a bacterial catalase-peroxidase enzyme KatG. Isoniazid is bactericidal to rapidly dividing mycobacteria and has anti-tuberculostatic activity.

 NH_2

99.68% Purity: Clinical Data: Launched

10 mM × 1 mL, 100 mg Size:

Isoniazid-d4 (INH-d4; Isonicotinic acid hydrazide-d4;

Isonicotinic hydrazide-d4) Cat. No.: HY-B0329S

Isoniazid-d4 (INH-d4) is the deuterium labeled Isoniazid. Isoniazid (INH) is a prodrug and must be activated by a bacterial catalase-peroxidase enzyme KatG. Isoniazid is bactericidal to rapidly dividing mycobacteria and has anti-tuberculostatic activity.

Purity: 98.95%

Clinical Data: No Development Reported Size 1 mg, 5 mg, 10 mg

Isorhapontigenin

Isorhapontigenin, an orally bioavailable dietary polyphenol isolated from the Chinese herb Gnetum cleistostachyum, displays anti-inflammatory effects. Isorhapontigenin induces autophagy and inhibits invasive bladder cancer formation.

99.82% Purity:

Clinical Data: No Development Reported 10 mM × 1 mL, 5 mg, 10 mg Size:



Cat. No.: HY-N2593

Isosorbide mononitrate

(Isosorbide-5-mononitrate) Cat. No.: HY-B0642

Isosorbide mononitrate(Isosorbide-5-mononitrate) is a nitrate-class compound used for angina pectoris; acts by dilating the blood vessels so as to reduce the blood pressure.

Purity: 99.89% Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g

Isotretinoin

(13-cis-Retinoic acid)

Isotretinoin(13-cis-Retinoic acid) is a medication used for the treatment of severe acne. It was first developed to be used as a chemotherapy medication for the treatment of brain cancer, pancreatic cancer and more.

Cat. No.: HY-15127

99.88% Clinical Data: Launched 100 mg, 500 mg

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Isotretinoin-d5

Cat. No.: HY-15127S

Isotretinoin-d5 (13-cis-Retinoic acid-d5) is the deuterium labeled Isotretinoin.

Isotretinoin(13-cis-Retinoic acid) is a medication used for the research of severe acne.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg

ISRIB (trans-isomer)

PERK with an IC₅₀ of 5 nM. ISRIB potently reverses the effects of eIF2α phosphorylation

Clinical Data: No Development Reported

Isradipine-d3

Isradipine-d3 (PN 200-110-d3) is the deuterium labeled Isradipine. Isradipine (PN 200-110) is an orally active L-type calcium channel blocker.

Cat. No.: HY-B0233S

Purity: >98%

Clinical Data

Size 1 mg, 10 mg

Itraconazole

(R51211) Cat. No.: HY-17514

Itraconazole (R51211) is a triazole antifungal agent and a potent and orally active Hedgehog (Hh) signaling pathway antagonist with an IC₅₀ of ~800 nM.



Purity: 99.15% Clinical Data: Launched Size: 100 mg, 500 mg

ITX5061

Cat. No.: HY-19900

ITX5061 is a type II inhibitor of p38 MAPK and also an antagonist of scavenger receptor B1 (SR-B1).



98.38% Purity:

Clinical Data: No Development Reported 5 mg, 10 mg, 50 mg

Size

Ivacaftor

(VX-770) Cat. No.: HY-13017

Ivacaftor (VX-770) is a potent and orally bioavailable CFTR potentiator, targeting G551D-CFTR and F508del-CFTR with ECsnS of 100 nM and 25 nM, respectively.



Purity: 99.90% Clinical Data: Launched

Size: $10 \text{ mM} \times 1 \text{ mL}, 5 \text{ mg}, 10 \text{ mg}$

Isradipine

(PN 200-110)

Isradipine (PN 200-110) is an orally active L-type calcium channel blocker. Isradipine, as a powerful peripheral vasodilator, is a dihydropyridine calcium antagonist with selective actions on the heart as well as the peripheral

Purity: 99 69% Clinical Data: Launched

10 mM × 1 mL, 10 mg, 50 mg, 100 mg Size:



Cat. No.: HY-12495

Cat. No.: HY-B0233

ISRIB (trans-isomer) is a potent inhibitor of

 $(IC_{50}=5 \text{ nM}).$

Purity: 99.37%

10 mM × 1 mL, 10 mg, 50 mg, 100 mg

Itraconazole-d5

Itraconazole-d5 (R51211-d5) is the deuterium labeled Itraconazole. Itraconazole (R51211) is a triazole antifungal agent and a potent and orally active Hedgehog (Hh) signaling pathway antagonist with an IC₅₀ of ~800 nM.

of the cooking

Cat. No.: HY-17514S

>98% Purity:

Clinical Data: No Development Reported

Size 500 μg, 1 mg

IU1

Cat. No.: HY-13817

IU1 is a special Usp14 inhibitor with an IC50 of 4-5 μΜ.



99.45% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg Size:

Ivacaftor benzenesulfonate

(VX-770 benzenesulfonate)

Cat. No.: HY-13017A

Ivacaftor benzenesulfonate is an orally bioavailable CFTR potentiator, used for cystic fibrosis treatment.



Purity: >98% Clinical Data: Launched 1 mg, 5 mg

Ivacaftor hydrate

(VX-770 hydrate) Cat. No.: HY-13017B

Ivacaftor hydrate (VX-770 hydrate) is an orally bioavailable CFTR potentiator, used for cystic fibrosis treatment.

Purity: >98% Clinical Data: Launched Size: 1 mg, 5 mg

Ivacaftor-d19

(VX-770-d19) Cat. No.: HY-13017S1

Ivacaftor-d19 (VX-770-d19) is the deuterium labeled Ivacaftor, Ivacaftor (VX-770) is a potent and orally bioavailable CFTR potentiator, targeting G551D-CFTR and F508del-CFTR with EC_{sn}s of 100 nM and 25 nM, respectively.



>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Ivermectin

(MK-933) Cat. No.: HY-15310

Ivermectin (MK-933) is a broad-spectrum anti-parasite agent. Ivermectin (MK-933) is a specific inhibitor of Impα/β1-mediated nuclear import and has potent antiviral activity towards both HIV-1 and dengue virus.



Purity: 96.79% Clinical Data: Launched

10 mM × 1 mL, 500 mg, 1 g Size:

Ixazomib

(MLN2238) Cat. No.: HY-10453

Ixazomib (MLN2238) is a selective, potent, and reversible proteasome inhibitor, which inhibits the chymotrypsin-like proteolytic (β5) site of the 20S proteasome with an IC_{50} of 3.4 nM (K_i of 0.93 nM).

Purity: 99.85% Clinical Data: Launched

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg

Ixazomib citrate

(MLN9708) Cat. No.: HY-10452

Ixazomib citrate (MLN9708) is a reversible inhibitor of the chymotrypsin-like proteolytic $\beta 5$ site of the 20S proteasome with an IC_{so} of 3.4 nM and a K, of 0.93 nM.



99.89% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

IZCZ-3

Cat. No.: HY-111411

IZCZ-3 is a potent c-MYC transcription inhibitor with antitumor activity.



99.45% Purity:

Clinical Data: No Development Reported Size 5 mg, 10 mg, 50 mg, 100 mg

JNJ-42165279

Cat. No.: HY-19636

JNJ-42165279 is a FAAH inhibitor with IC50 of 70 \pm 8 nM and 313 \pm 28 nM for hFAAH and rFAAH, respectively.

99.87% Purity: Clinical Data: Phase 2

10 mM × 1 mL, 5 mg, 10 mg, 25 mg Size:

Juglanin

Cat. No.: HY-N3442

Juglanin, a natural occurring flavonoid, is a JNK acticator, with inflammation and anti-tumor activities. Juglanin can induce apoptosis and autophagy on human breast cancer cells.



99.90% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

JZL195

Cat. No.: HY-15250

JZL195 is a selective and efficacious dual fatty acid amide hydrolase (FAAH) and monoacylglycerol lipase (MAGL) inhibitor with IC₅₀s of 2 and 4 nM, respectively.



Purity: 99.81%

Clinical Data: No Development Reported 5 mg, 10 mg, 25 mg, 50 mg, 100 mg Size:

K-252a

(SF2370; Antibiotic K 252a; Antibiotic SF 2370)

K-252a, a staurosporine analog, inhibits protein kinase, with IC₅₀ values of 470 nM, 140 nM, 270 nM, and 1.7 nM for PKC, PKA, Ca²⁺/calmodulin-dependent kinase type II, and

phosphorylase kinase, respectively.

Purity: 99.45%

Clinical Data: No Development Reported $10 \text{ mM} \times 1 \text{ mL}, 1 \text{ mg}, 5 \text{ mg}$



Cat. No.: HY-N6732

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

Kaempferide

(Kaempferol 4'-O-methyl ether)

Cat. No.: HY-15449

Kaempferide is an O-methylated flavonol, a type of chemical compound. It can be found in Kaempferia galanga (aromatic ginger).

Purity: 99 42%

Clinical Data: No Development Reported Size: $10 \text{ mM} \times 1 \text{ mL}$, 5 mg, 10 mg, 50 mg Kaempferol

(Kempferol; Robigenin)

Kaempferol (Kempferol), a flavonoid found in many edible plants, inhibits estrogen receptor $\boldsymbol{\alpha}$ expression in breast cancer cells and induces apoptosis in glioblastoma cells and lung cancer cells by activation of MEK-MAPK. Kaempferol can be uesd for the research of breast cancer.



Cat. No.: HY-15415

Cat. No.: HY-14590

99.67% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 50 mg, 100 mg, 200 mg, 500 mg

KAN0438757

Cat. No.: HY-112808

KAN0438757 is a potent and selective inhibitor of the metabolic kinase PFKFB3 with an IC_{50} of 0.19 μΜ.

Purity: 99 30%

Clinical Data: No Development Reported 10 mM × 1 mL, 5 mg, 10 mg, 50 mg

KB-R7943 mesylate

KB-R7943 mesylate is a widely used inhibitor of the reverse Na+/Ca2+ exchanger (NCX_{rev}) with IC_{50} of 5.7±2.1 µM. KB-R7943 mesylate induces

cancer cell death via activating the JNK pathway and blocking autophagic flux.

Purity: 99.16%

Clinical Data: No Development Reported 10 mM × 1 mL, 10 mg, 50 mg

Ketanserin tartrate

(R41468 tartrate) Cat. No.: HY-10562A

Ketanserin (R41468) tartrate is a selective 5-HT2

receptor antagonist. Ketanserin tartrate also blocks hERG current ($I_{\rm hERG}$) in a concentration-dependent manner (IC₅₀=0.11 μM).



99.99% Purity: Clinical Data: Launched

10 mM \times 1 mL, 50 mg, 100 mg

Ketanserin

(R41468) Cat. No.: HY-10562

Ketanserin is a selective 5-HT2 receptor antagonist. Ketanserin also blocks hERG current (I_{hERG}) in a concentration-dependent manner $(IC_{50}=0.11 \mu M).$

99.24% Purity: Clinical Data: Launched

10 mM × 1 mL, 50 mg, 100 mg Size:

KJ Pyr 9

Cat. No.: HY-19735

KJ Pyr 9 is an inhibitor of MYC with a K_d of 6.5 nM in in vitro assay.

Purity: 99.29%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

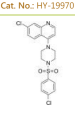
KM11060

KM11060 is a corrector of the F508 deletion (F508del)-cystic fibrosis transmembrane conductance regulator (CFTR) trafficking defect. KM11060 can be used for the research of F508del-CFTR processing defect and development of

cystic fibrosis therapeutics. Purity: 99.59%

Clinical Data: No Development Reported

10 mM × 1 mL, 10 mg, 25 mg, 50 mg, 100 mg



KN-93

Cat. No.: HY-15465

KN-93 is a cell-permeable, reversible and competitive inhibitor calmodulin-dependent kinase type II (CaMKII) with a K, of 370 nM.



Purity: 99.19%

No Development Reported Clinical Data:

Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 25 mg, 50 mg

KN-93 hydrochloride

Cat. No.: HY-15465A

KN-93 hydrochloride is a cell-permeable, reversible and competitive inhibitor calmodulin-dependent kinase type II (CaMKII) with a K_i of 370 nM.



99.92%

Clinical Data: No Development Reported

10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 25 mg, 50 mg

KN-93 phosphate

KN-93 phosphate is a novel membrane-permeant synthetic inhibitor of purified neuronal CaMK-II, with K, of 370 nM.



Cat. No.: HY-15465B

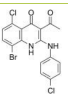
Purity: 99.69%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 50 mg

KSI-3716

KSI-3716 is a potent **c-Myc** inhibitor that blocks c-MYC/MAX binding to target gene promoters. KSI-3716 is an effective intravesical chemotherapy agent for bladder cancer.



Cat. No.: HY-12703

Purity: 98.55%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

KU-55933

Cat. No.: HY-12016

KU-55933 is a potent **ATM** inhibitor with an IC_{50} and K_i of 12.9 and 2.2 nM, respectively, and is highly selective for ATM as compared to DNA-PK, PJ3K/PJ4K, ATR and mTOR.



Purity: 99.88%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

L-779450

Cat. No.: HY-12787

L-779450 is a potent and selective **B-Raf** kinase inhibitor with a $K_{\rm d}$ of 2.4 nM.

N Co

Purity: 98.88%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Laduviglusib

(CHIR-99021; CT99021) Cat. No.: HY-10182

Laduviglusib (CHIR-99021) is a potent and selective $GSK-3\alpha/\beta$ inhibitor with IC_{50} s of 10 nM and 6.7 nM. Laduviglusib shows >500-fold selectivity for GSK-3 over CDC2, ERK2 and other protein kinases.

Purity: 99.76%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Laduviglusib monohydrochloride

(CHIR-99021 monohydrochloride; CT99021 monohydrochloride)t. No.: HY-10182A

Laduviglusib (CHIR-99021) monohydrochloride is a potent and selective GSK-3 α / β inhibitor with IC₅₀S of 10 nM and 6.7 nM. Laduviglusib monohydrochloride shows >500-fold selectivity for GSK-3 over CDC2, ERK2 and other protein kinases.



Purity: 99.93%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Laduviglusib trihydrochloride

(CHIR-99021 trihydrochloride; CT99021 trihydrochloride) Cat. No.: HY-10182B

Laduviglusib (CHIR-99021) trihydrochloride is a potent and selective $GSK\text{-}3\alpha/\beta$ inhibitor with $IC_{sg}\text{S}$ of 10 nM and 6.7 nM. Laduviglusib trihydrochloride shows >500-fold selectivity for GSK-3 over CDC2, ERK2 and other protein kinases.



Purity: 98.68%

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

Lamotrigine

(LTG; BW430C)

Lamotrigine (BW430C) is a potent and orally active anticonvulsant or antiepileptic agent. Lamotrigine selectively blocks voltage-gated Na* channels, stabilizing presynaptic neuronal membranes and inhibiting glutamate release.



Cat. No.: HY-B0495

Purity: 99.86% Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g

Lamotrigine-13C,d3

(LTG-13C,d3; BW430C-13C,d3)

Cat. No.: HY-B0495S1

Lamotrigine-13C,d3 is the 13C- and deuterium labeled. Lamotrigine (BW430C) is a potent and orally active anticonvulsant or antiepileptic agent.

Purity: > 98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Lamotrigine-13C3,d3

(LTG-13C3,d3; BW430C-13C3,d3)

Lamotrigine-13C3,d3 (LTG-13C3,d3) is the 13C-labeled Lamotrigine. Lamotrigine (BW430C) is a potent and orally active **anticonvulsant** or antiepileptic agent.



Cat. No.: HY-B0495S

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

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Lanatoside C

Cat. No.: HY-B1030

Lanatoside C is a cardiac glycoside, can be used in the treatment of congestive heart failure and cardiac arrhythmia.Lanatoside C has an IC50 of $0.19 \mu M$ for dengue virus infection in HuH-7 cells.



Purity: 99 81% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg

Lapatinib

(GW572016; GW2016)

Lapatinib (GW572016) is a potent inhibitor of the ErbB-2 and EGFR tyrosine kinase domains with IC_{so} values against purified EGFR and ErbB-2 of 10.2 and 9.8 nM, respectively.



Cat. No.: HY-50898

Purity: 99.83% Clinical Data: Launched

Size: 10 mM × 1 mL, 50 mg, 100 mg, 500 mg, 1 g

Lapatinib ditosylate (GW572016 ditosylate monohydrate; GW2016

ditosylate monohydrate)

Lapatinib ditosylate monohydrate (GW572016 ditosylate monohydrate) is a potent inhibitor of the ErbB-2 and EGFR tyrosine kinase domains with IC₅₀ values against purified EGFR and ErbB-2 of 10.2 and 9.8 nM, respectively.



Cat. No.: HY-50898B

Purity: 99 78% Clinical Data: Launched

10 mM × 1 mL, 50 mg, 100 mg Size:

Lapatinib ditosylate

(GW572016 ditosylate; GW2016 ditosylate)

Lapatinib ditosylate (GW572016 ditosylate) is a potent inhibitor of the ErbB-2 and EGFR tyrosine kinase domains with IC₅₀ values against purified EGFR and ErbB-2 of 10.2 and 9.8 nM, respectively.



Cat. No.: HY-50898A

Purity: 99 95% Clinical Data: Launched

10 mM × 1 mL, 50 mg, 100 mg, 500 mg, 1 g

Lapatinib-d4-1

(GW572016-d4-1; GW2016-d4-1)

Lapatinib-d4-1 is deuterium labeled Lapatinib. Lapatinib (GW572016) is a potent inhibitor of the ErbB-2 and EGFR tyrosine kinase domains with IC50 values against purified EGFR and ErbB-2 of 10.2 and 9.8 nM, respectively.



Cat. No.: HY-50898S3

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Lapatinib-d5

(GW572016-d5; GW2016-d5)

Lapatinib-d5 is deuterium labeled Lapatinib. Lapatinib (GW572016) is a potent inhibitor of the ErbB-2 and EGFR tyrosine kinase domains with IC50 values against purified EGFR and ErbB-2 of 10.2 and 9.8 nM, respectively.



Cat. No.: HY-50898S2

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Lapatinib-d7 dihydrochloride

(GW572016-d7 dihydrochloride; GW2016-d7 dihydrochloridetat. No.: HY-50898S1

Lapatinib-d7 (GW572016-d7) dihydrochloride is the deuterium labeled Lapatinib dihydrochloride. Lapatinib (GW572016) dihydrochloride is a potent inhibitor of the ErbB-2 and EGFR tyrosine kinase domains with IC_{50} values against purified EGFR and ErbB-2 of 10.2 and 9.8 nM, respectively.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 ma. 5 ma

Lapatinib-d7 ditosylate

Cat. No.: HY-50898BS

Lapatinib-d7 (GW572016-d7) ditosylate is the deuterium labeled Lapatinib. Lapatinib (GW572016) is a potent inhibitor of the ErbB-2 and EGFR tyrosine kinase domains with IC_{so} values against purified EGFR and ErbB-2 of 10.2 and 9.8 nM, respectively.



Size: 1 mg, 10 mg



Lasalocid

(Lasalocid-A; Ionophore X-537A; Antibiotic X-537A) Cat. No.: HY-B1071

Lasalocid (Lasalocid-A; Ionophore X-537A; Antibiotic X-537A) is an antibacterial agent and a coccidiostat, used in the feed additives.



Purity: 96.33%

Clinical Data: No Development Reported

Size 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Lasalocid sodium (Lasalocid-A sodium; Ionophore X-537A Cat. No.: HY-B1071A

sodium; Antibiotic X-537A sodium)

Lasalocid sodium (Lasalocid-A sodium) treatment led to an increase in cell wall thickness, whilst the quantity and sugar composition of the cell wall remained unchanged in BY-2 cells.



Purity: ≥97.0%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Latrepirdine dihydrochloride

(Dimebolin dihydrochloride)

Cat. No.: HY-14537

Latrepirdine dihydrochloride is a neuroactive compound with antagonist activity at histaminergic, α -adrenergic, and serotonergic receptors. Latrepirdine stimulates amyloid precursor protein (APP) catabolism and amyloid- β (A β) secretion.

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Purity: 99.75% Clinical Data: Launched

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg, 200 mg

LC3-mHTT-IN-AN1

LC3-mHTT-IN-AN1 (Compound AN1) is a mHTT-LC3 linker compound, which interacts with both mutant huntingtin protein (mHTT) and LC3B but not with wtHTT or irrelevant control proteins.



Cat. No.: HY-130258

Purity: 97.14%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

LC3-mHTT-IN-AN2

Cat. No.: HY-130259

LC3-mHTT-IN-AN2 (Compound AN2) is a mHTT-LC3 linker compound, which interacts with both mutant huntingtin protein (mHTT) and LC3B but not with wtHTT or irrelevant control proteins.



Purity: 96.07%

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

Leonurine

(SCM-198) Cat. No.: HY-N0741

Leonurine is an alkaloid isolated from Herba leonuri, with anti-oxidative and anti-inflammatory.



Purity: 99.62%

Clinical Data: No Development Reported

Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Leonurine hydrochloride

(SCM-198 hydrochloride) Cat. No.: HY-N0741A

Leonurine hydrochloride is an alkaloid isolated from Herba leonuri, with anti-oxidative and anti-inflammatory.

Purity: 99.69%

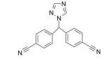
Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Letrozole

(CGS 20267) Cat. No.: HY-14248

Letrozole (CGS 20267) is a potent, selective, reversible and orally active non-steroidal inhibitor of aromatase, with an $\rm IC_{50}$ of 11.5 nM. Letrozole selective inhibits estrogen biosynthesis, and can be used for the research of breast cancer.



Purity: 99.92% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 200 mg, 500 mg

Letrozole-d4

Cat. No.: HY-14248S

Letrozole-d4 (CGS 20267-d4) is the deuterium labeled Letrozole. Letrozole (CGS 20267) is a potent, selective, reversible and orally active non-steroidal inhibitor of **aromatase**, with an $\rm IC_{50}$ of 11.5 nM.



Purity: >98%

Clinical Data:

Size: 1 mg, 10 mg

Levosimendan

(Simsndan; OR-1259)

Levosimendan (Simsndan; OR-1259) is a calcium sensitiser used in the management of acutely decompensated congestive heart failure.



Cat. No.: HY-14286

Purity: 99.51% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg

Levosimendan D3

(Simsndan D3; OR-1259 D3) Cat. No.: HY-14286S

Levosimendan D3 (Simsndan D3) is a deuterium labeled Levosimendan. Levosimendan is a calcium sensitiser used in the management of acutely decompensated congestive heart failure.



Purity: > 98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

LG-100064

Cat. No.: HY-104070

LG-100064 is a **retinoid-X-receptor (RXR)** agonist, with EC_{so} s of 330 nM, 200 nM, and 260 nM for RXR α , RXR β and RXR γ ; LG-100064 can be used in the research of cancer.



Purity: 99.52%

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 5 mg, 10 mg

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

LG100268

(LG268) Cat. No.: HY-15340

LG100268 (LG268) is a potent, selective and orally active retinoid X receptor (RXR) agonist with EC_{50} values of 4 nM, 3 nM, and 4 nM for RXR- α , RXR-β, and RXR-γ, respectively.



Purity: 99 22%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Licochalcone E

Licochalcone E, a flavonoid compound isolated from Glycyrrhiza inflate, inhibits NF- κB and AP-1 transcriptional activity through the inhibition of AKT and MAPK activation.

Cat. No.: HY-N4182

Purity: 99 63%

Clinical Data: No Development Reported

5 mg, 10 mg

Liensinine Diperchlorate

Cat. No.: HY-N0485

Liensinine Diperchlorate is a major isoquinoline alkaloid, extracted from the seed embryo of Nelumbo nucifera Gaertn. Liensinine Diperchlorate inhibits late-stage autophagy/mitophagy through blocking autophagosome-lysosome fusion.



Clinical Data: No Development Reported Size: 5 mg, 10 mg, 20 mg



Linagliptin

(BI 1356) Cat. No.: HY-10284

Linagliptin is a highly potent, selective DPP-4 inhibitor with IC₅₀ of 1 nM.



Purity: 99.97% Clinical Data: Launched

10 mM × 1 mL, 100 mg, 500 mg, 1 g Size:

Linagliptin-d4

(BI 1356-d4) Cat. No.: HY-10284S

Linagliptin-d4 is deuterium labeled Linagliptin. Linagliptin is a highly potent, selective DPP-4 inhibitor with IC50 of 1 nM.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Licochalcone A

(Licochalcone-A) Cat. No.: HY-N0372

Licochalcone A (LCA), a flavonoid isolated, presents obvious anti-cancer effects, displays broad-spectrum inhibition against UDP-glucuronosyltransferases (UGTs).



Purity: 99 89% Clinical Data: Phase 1

Size: 5 mg, 10 mg, 50 mg, 100 mg

Liensinine

Cat. No.: HY-N0484

Liensinine is an autophagy/mitophagy inhibitor.



Cat. No.: HY-N0401

Purity: 99 89%

Clinical Data: No Development Reported 5 mg, 10 mg, 20 mg

Ligustilide

Ligustilide is is a bioactive phthalide derivative isolated from Angelica sinensis and Chuanxiong. Ligustilide exhibits neuroprotective, anti-cancer, anti-inflammatory, and vasodilator effects.



Purity: 98.49%

Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg

Linagliptin-13C,d3

(BI 1356-13C,d3) Cat. No.: HY-10284S1

Linagliptin-13C,d3 is the 13C- and deuterium labeled. Linagliptin is a highly potent, selective DPP-4 inhibitor with IC50 of 1 nM.



Purity: >98%

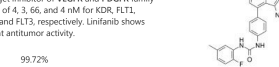
Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Linifanib

(ABT-869; AL-39324)

Linifanib (ABT-869) is a potent and orally active multi-target inhibitor of VEGFR and PDGFR family with IC_{so}s of 4, 3, 66, and 4 nM for KDR, FLT1, PDGFRβ, and FLT3, respectively. Linifanib shows prominent antitumor activity.



Purity: Clinical Data: Phase 3

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

Cat. No.: HY-50751

Lithocholic acid

(3α-Hydroxy-5β-cholanic acid)

Lithocholic acid is a toxic secondary bile acid, causes intrahepatic cholestasis, has tumor-promoting activity. Target: Others Lithocholic acid has been used in a study to assess cholestasis and its action on several organs and tissues in rats.

HO H H H OH

Cat. No.: HY-B0172

Purity: ≥98.0%

Lithocholic acid-d5

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g

Lithocholic acid-d4

(3α-Hydroxy-5β-cholanic acid-d4)

Lithocholic acid-d4 (3 α -Hydroxy-5 β -cholanic acid-d4) is the deuterium labeled Lithocholic acid, which is a toxic secondary bile acid.



Cat. No.: HY-B0172S

Purity: >98%

Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg, 25 mg

(3α-Hydroxy-5β-cholanic acid-d5)

Lithocholic acid-d5 is deuterium labeled Lithocholic acid



Cat. No.: HY-B0172S1

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Lomustine

(CCNU; NSC 79037)

Lomustine (CCNU; NSC 79037) is a **DNA alkylating** agent, with antitumor activity.



Cat. No.: HY-13669

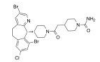
Purity: 99.91% Clinical Data: Launched

Size: 10 mM × 1 mL, 200 mg, 500 mg

Lonafarnib

(Sch66336) Cat. No.: HY-15136

Lonafarnib (Sch66336) is a potent and orally active farnesyl transferase (FTase) inhibitor. Lonafarnib inhibits the activities of H-ras, K-ras and N-ras with IC₅₀ values of 1.9 nM, 5.2 nM and 2.8 nM, respectively. Lonafarnib also has anti-hepatitis delta virus (HDV) activities.



Purity: 98.67% Clinical Data: Launched

Size: $10 \text{ mM} \times 1 \text{ mL}, 5 \text{ mg}, 10 \text{ mg}$

Loperamide hydrochloride

(R-18553 hydrochloride)

Loperamide (hydrochloride) (R-18553 (hydrochloride)) is an **opioid receptor** agonist. Loperamide hydrochloride is a selective and competitive human intestinal carboxylesterases (hiCE) inhibitor. Loperamide hydrochloride has anti-diarrheal effect.

Purity: 99.79% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg



Cat. No.: HY-B0418A

Loperamide-d6 hydrochloride

(R-18553-d6 hydrochloride)

Loperamide D6 hydrochloride (R-18553 D6 hydrochloride) is a deuterium labeled Loperamide hydrochloride. Loperamide hydrochloride is an **opioid receptor** agonist for the treatment of diarrhea.



Cat. No.: HY-B0418AS

Purity: > 98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Loperamide-d6 N-Oxide

Loperamide-d6 N-Oxide is the deuterium labeled Loperamide hydrochloride. Loperamide hydrochloride (R-18553 hydrochloride) is an **opioid receptor** agonist. Loperamide hydrochloride is a selective and competitive human intestinal carboxylesterases

(hiCE) inhibitor.

Purity: >98%

Clinical Data: No Development Reported Size: 2.5 mg, 1 mg, 5 mg, 10 mg



Cat. No.: HY-B0418AS1

Losmapimod

(GSK-AHAB; GW856553X; SB856553)

Losmapimod (GSK-AHAB) is a selective, potent, and orally active **p38 MAPK** inhibitor with **pK**₁s of 8.1 and 7.6 for p38 α and p38 β , respectively.



Cat. No.: HY-10402

Purity: 98.06% Clinical Data: Phase 3

Size: 10 mM × 1 mL, 10 mg, 50 mg

Lovastatin (Mevinolin)

Lovastatin is a cell-permeable HMG-CoA reductase inhibitor used to lower cholesterol.



Cat. No.: HY-N0504

Purity: 99.93% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg

Lovastatin-d3

(Mevinolin-d3) Cat. No.: HY-N0504S2

Lovastatin-d3 is deuterium labeled Lovastatin. Lovastatin is a cell-permeable HMG-CoA reductase inhibitor used to lower cholesterol.

Purity: > 98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Lovastatin-d9

Lovastatin-d9 is the deuterium labeled Lovastatin. Lovastatin is a cell-permeable HMG-CoA reductase inhibitor used to lower cholesterol.



Cat. No.: HY-N0504S1

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Lucanthone

Cat. No.: HY-B2098

Lucanthone is an endonuclease inhibitor of Apurinic endonuclease-1 (APE-1).

Purity: 99.71% Clinical Data: Phase 2

Size: 1 mg, 5 mg, 10 mg, 25 mg

Lucanthone-d4 hydrochloride

Cat. No.: HY-B2098AS

Lucanthone-d4 hydrochloride is the deuterium labeled Lucanthone. Lucanthone is an endonuclease inhibitor of Apurinic endonuclease-1 (APE-1).



Purity: >98% Clinical Data:

Size: 1 mg, 10 mg

Lumacaftor

(VX-809; VRT 826809) Cat. No.: HY-13262

Lumacaftor (VX-809; VRT 826809) is a CFTR modulator that corrects the folding and trafficking of CFTR protein.



Purity: 99.19% Clinical Data: Launched

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Lumefantrine

(Benflumetol) Cat. No.: HY-B0803

Lumefantrine is an antimalarial drug, used in combination with Artemether. The artemether-lumefantrine (AL) as the first- and second-line anti-malarial drugs.



Purity: 98.41% Clinical Data: Launched

Size: 10 mg, 50 mg, 100 mg, 500 mg

Lumefantrine-d9

(Benflumetol-d9) Cat. No.: HY-B0803S1

Lumefantrine-d9 (Benflumetol-d9) is the deuterium labeled Lumefantrine. Lumefantrine is an antimalarial drug, used in combination with Artemether. The artemether-lumefantrine (AL) as the first- and second-line anti-malarial drugs.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Luminespib

(VER-52296; AUY922; NVP-AUY922) Cat. No.: HY-10215

Luminespib (VER-52296) is a potent HSP90 inhibitor with IC_{so} s of 7.8 and 21 nM for HSP90 α and HSP90 β , respectively.



Purity: 99.89%

Clinical Data: No Development Reported

Size: $10 \text{ mM} \times 1 \text{ mL}$, 5 mg, 10 mg, 25 mg, 100 mg, 200 mg

Luteolin

(Luteoline; Luteolol; Digitoflavone) Cat. No.: HY-N0162

Luteolin (Luteoline), a flavanoid compound, is a potent Nrf2 inhibitor.



Purity: 98.42% Clinical Data: Phase 1

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 500 mg

LV-320

LV-320 is a potent and uncompetitive ATG4B inhibitor with an IC $_{50}$ of 24.5 μ M and a K $_{d}$ of 16 μ M. LV-320 inhibits ATG4B enzymatic activity, blocks autophagic flux in cells, and is stable,

non-toxic and active in vivo.

Purity: ≥95.0%

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



Cat. No.: HY-112711

LX2343

LX2343 is a BACE1 enzyme inhibitor with an IC_{so} value of 11.43 \pm 0.36 μ M. LX2343 acts as a non-ATP competitive PI3K inhibitor with an IC₅₀ of $15.99\pm3.23~\mu\text{M}$. LX2343 stimulates autophagy in its

promotion of $A\beta$ clearance.

Cat. No.: HY-111383

99 80% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

LY2109761

LY2109761 is an orally active, selective TGF-β receptor type I/II inhibitor with K.s of 38 nM and 300 nM, respectively.



Cat. No.: HY-12075

99 88% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

LY2183240

Cat. No.: HY-10865

LY2183240 is a highly potent blocker of anandamide uptake (IC_{s0}= 270 pM; K_i=540 nM). LY2183240 is a potent, covalent inhibitor of the endocannabinoid-degrading enzyme fatty acid amide hydrolase (FAAH) with an IC_{so} of 12.4 nM.

Purity: 99.07%

Clinical Data: No Development Reported 10 mM × 1 mL, 10 mg, 50 mg

LY2562175

Cat. No.: HY-103704

LY2562175 is a potent and selective FXR agonist,

with an EC_{50} of 193 nM.



99.26%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg

LY294002

Cat. No.: HY-10108

LY294002 is a broad-spectrum inhibitor of PI3K with $IC_{50}s$ of 0.5, 0.57, and 0.97 μM for $PI3K\alpha$, $PI3K\delta$ and PI3Kβ, respectively. LY294002 also inhibits CK2 with an IC₅₀ of 98 nM.



99.95% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg, 500 mg

LY2955303

Cat. No.: HY-107765

LY2955303 is a potent and selective retinoic acid receptor gamma (RARγ) antagonist with a K_i of 1.09 nM.



Purity: 99.16%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg

LY3009120

(DP-4978) Cat. No.: HY-12558

LY3009120 (DP-4978) is a pan RAF inhibitor which inhibits BRAFV600E, BRAFWT and CRAFWT with IC_{so}s of 5.8, 9.1 and 15 nM, respectively.



Purity: 99.01% Clinical Data: Phase 1

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Lycorine hydrochloride

Lycorine hydrochloride is the main active ingredient of the herbal medicine derived from Lycoris radia and is also a melanoma vasculogenic inhibitor and has anti-tumor activity. Lycorine hydrochloride effectively inhibits mitotic proliferation of Hey1B cells (IC $_{50}$ of 1.2 μ M).



Cat. No.: HY-N0289

Purity: 99.89%

Clinical Data: No Development Reported

10 mM × 1 mL, 10 mg, 25 mg, 50 mg, 100 mg

LYN-1604

Cat. No.: HY-101923

LYN-1604 is a potent UNC-51-like kinase 1 (ULK1) activator (EC_{50} =18.94 nM) for the research of triple negative breast cancer (TNBC).



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

LYN-1604 dihydrochloride

Cat. No.: HY-101923B

LYN-1604 dihydrochloride is a potent UNC-51-like kinase 1 (ULK1) activator (EC₅₀=18.94 nM) for the research of triple negative breast cancer (TNBC).



98.73%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

LYN-1604 hydrochloride

Cat. No.: HY-101923A

LYN-1604 hydrochloride 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%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Lys05

(Lys01 trihydrochloride)

Lys05 (Lys01 trihydrochloride) is a novel lysosomal autophagy inhibitor with IC_{so} values of 3.6, 3.8, 6 and 7.9 μM for 1205Lu, c8161, LN229 and HT-29 cell line in the MTT assay.



Cat. No.: HY-12855A

99 39% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Magnolol

Cat. No.: HY-N0163

Magnolol, a natural lignan isolated from the stem bark of Magnolia officinalis, is a dual agonist of both RXR α and PPAR γ , with EC₅₀ values of 10.4 μM and 17.7 μM, respectively.

Purity: 99 92% Clinical Data: Launched

10 mM × 1 mL, 10 mg, 50 mg, 100 mg

Malvidin-3-O-arabinoside chloride

Cat. No.: HY-N9349

Malvidin-3-O-arabinoside chloride ameliorates ethyl carbamate-induced oxidative damage by stimulating AMPK-mediated autophagy.



Purity: >98%

Clinical Data: No Development Reported 5 mg, 10 mg, 25 mg

Manzamine A hydrochloride

Cat. No.: HY-117025A

Manzamine A hydrochloride, an orally active beta-carboline alkaloid, inhibits specifically GSK-3 β and CDK-5 with IC_{sn}s of 10.2 μ M and 1.5 μM, respectively. Manzamine A hydrochloride targets vacuolar ATPases and inhibits autophagy in pancreatic cancer cells.

Purity: 99 29%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

MAPK13-IN-1

Cat. No.: HY-18850

MPAK13-IN-1 is a MAPK13 (p38 δ) inhibitor, with an IC₅₀ of 620 nM.



99.63% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Maprotiline hydrochloride

Cat. No.: HY-B0444

Maprotiline hydrochloride is a selective noradrenalin re-uptake inhibitor and a tetracyclic antidepressant. Target: Others Maprotiline (sold as Deprilept, Ludiomil, Psymion) is a tetracyclic antidepressant (TeCA).



HCI

99.43% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg

Maprotiline-d3 hydrochloride

Cat. No.: HY-B0444S1

Maprotiline-d3 (hydrochloride) is deuterium labeled Maprotiline (hydrochloride).

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Maprotiline-d5 hydrochloride

Cat. No.: HY-B0444S

Maprotiline-d5 hydrochloride is the deuterium labeled Maprotiline hydrochloride. Maprotiline hydrochloride is a selective noradrenalin re-uptake inhibitor and a tetracyclic antidepressant.



Purity: >98%

Clinical Data:

Size: 1 mg, 10 mg

Matrine

(Matridin-15-one; Vegard; α-Matrine)

Matrine (Matridin-15-one) is an alkaloid found in plants from the Sophora genus. It has a variety of pharmacological effects, including anti-cancer effects, and action as a kappa opioid receptor and u-receptor agonist.



10 mM × 1 mL, 100 mg, 200 mg, 500 mg



Cat. No.: HY-N0164

Mdivi-1

(Mitochondrial division inhibitor 1)

Cat. No.: HY-15886

Mdivi-1 is a selective dynamin-related protein 1 (Drp1) inhibitor, Mdivi-1 is a mitochondrial division/mitophagy inhibitor.

Purity: 99 73%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Mefloquine hydrochloride

(Mefloquin hydrochloride)

Mefloquine hydrochloride (Mefloquin hydrochloride), a quinoline antimalarial agent, is an anti-SARS-CoV-2 entry inhibitor. Mefloquine hydrochloride is also a K+ channel

(KvQT1/minK) antagonist with an IC_{so} of \sim 1 μ M.

99 98% Purity: Clinical Data: Launched

10 mM × 1 mL, 100 mg, 500 mg



Cat. No.: HY-17437A

Size:

Megestrol acetate

Cat. No.: HY-13676

Megestrol acetate is a synthetic and orally active progesteronal agent. Megestrol acetate is effective as an appetite stimulant for wasting syndromes such as cachexia. Megestrol acetate decreases nuclear and cytosol androgen receptors human BPH tissue.



Purity: Clinical Data: Launched

10 mM × 1 mL, 500 mg, 1 g, 5 g Size:

Megestrol acetate-d3

Cat. No.: HY-13676S

Megestrol acetate-d3 is the deuterium labeled Megestrol acetate. Megestrol acetate is a synthetic and orally active progesteronal agent. Megestrol acetate is effective as an appetite stimulant for wasting syndromes such as cachexia.

Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg

Megestrol acetate-d3-1

Cat. No.: HY-13676S1

Megestrol acetate-d3-1 is deuterium labeled Megestrol acetate. Megestrol acetate is a synthetic and orally active progesteronal agent. Megestrol acetate is effective as an appetite stimulant for wasting syndromes such as cachexia.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Meglutol

(Dicrotalic acid; 3-Hydroxy-3-methylglutaric acid) Cat. No.: HY-B1189

Meglutol is an antilipemic agent which lowers cholesterol, triglycerides, serum beta-lipoproteins and phospholipids, and inhibits the activity of hydroxymethylglutarryl CoA reductases, which is the rate limiting enzyme in the biosynthesis of cholesterol.

≥98.0% **Purity:**

Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg

Melatonin

(N-Acetyl-5-methoxytryptamine) Cat. No.: HY-B0075

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.



99.73% Purity: Clinical Data: Launched

10 mM × 1 mL, 500 mg, 1 g, 5 g Size:

Melatonin-d3

(N-Acetyl-5-methoxytryptamine-d3)

Melatonin-d3 (N-Acetyl-5-methoxytryptamine-d3) is the deuterium labeled Melatonin. Melatonin is a hormone made by the pineal gland that can activates melatonin receptor.

Cat. No.: HY-B0075S1

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Melatonin-d4

(N-Acetyl-5-methoxytryptamine-d4) Cat. No.: HY-B0075S

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.



Purity: 95.87%

Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg

Melatonin-d7

(N-Acetyl-5-methoxytryptamine-d7)

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.



Cat. No.: HY-B0075S2

Purity: >98%

Clinical Data: No Development Reported 1 mg, 5 mg, 10 mg

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

Meloxicam

Cat. No.: HY-B0261

Meloxicam is a non-steroidal antiinflammatory agent, inhibits COX activity, with IC_{so} of 0.49 μM and 36.6 μM for COX-2 and COX-1, respectively.

Purity: 99.88% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg

Meloxicam-13C,d3

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.



Cat. No.: HY-B0261S2

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Meloxicam-d3

Cat. No.: HY-B0261S

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.

Purity: >98%

Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg

Meloxicam-d3-1

Cat. No.: HY-B0261S1

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.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Meprednisone

Cat. No.: HY-B0243

Meprednisone is a glucocorticoid and a methylated derivative of prednisone. Target: Glucocorticoid Receptor Meprednisone is a glucocorticoid and a methylated derivative of prednisone.



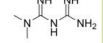
Purity: 99.60% Clinical Data: Launched

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Metformin

(1,1-Dimethylbiquanide)

Metformin (1,1-Dimethylbiguanide) inhibits the mitochondrial respiratory chain in the liver, leading to activation of AMPK, enhancing insulin sensitivity for type 2 diabetes research. Metformin can cross the blood-brain barrier and triggers autophagy.



Cat. No.: HY-B0627

Purity: 99.64% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 25 mg

Metformin hydrochloride

(1,1-Dimethylbiguanide hydrochloride)

Metformin hydrochloride (1,1-Dimethylbiguanide hydrochloride) inhibits the **mitochondrial respiratory chain** in the liver, leading to activation of AMPK, enhancing insulin sensitivity for type 2 diabetes research. Metformin hydrochloride triggers **autophagy**.

NH NH N NH₂

Cat. No.: HY-17471A

Purity: 99.89% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 25 mg

Metformin-d6 hydrochloride

(1,1-Dimethylbiguanide-d6 hydrochloride)

Metformin D6 hydrochloride is a deuterium labeled Metformin hydrochloride. Metformin hydrochloride inhibits the mitochondrial respiratory chain in the liver, leading to activation of AMPK, enhancing insulin sensitivity for type 2 diabetes research.



Cat. No.: HY-110228

Purity: ≥98.0%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Methylprednisolone-d2

(U 7532-d2) Cat. No.: HY-B0260S4

Methylprednisolone-d2 is the deuterium labeled Methylprednisolone. Methylprednisolone is a synthetic corticosteroid with anti-inflammatory and immunomodulating properties.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Methylprednisolone

(U 7532) Cat. No.: HY-B0260

Methylprednisolone is a synthetic corticosteroid with anti-inflammatory and immunomodulating properties. Methylprednisolone improve severe or critical COVID-19 by activating ACE2 and reducing IL-6 levels.



Purity: 99.75%
Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg

Methylprednisolone-d3

(U 7532-d3) Cat. No.: HY-B0260S

Methylprednisolone-d3 (U 7532-d3) is the deuterium labeled Methylprednisolone. Methylprednisolone is a synthetic corticosteroid with anti-inflammatory and immunomodulating properties.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Methyl prednisolone-d 4

(U 7532-d4) Cat. No.: HY-B0260S2

Methylprednisolone-d4 is deuterium labeled Methylprednisolone. Methylprednisolone is a synthetic corticosteroid with anti-inflammatory and immunomodulating properties.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Methylprednisolone-d5

(U 7532-d5) Cat. No.: HY-B0260S1

Methylprednisolone-d5 (U 7532-d5) is the deuterium labeled Methylprednisolone. Methylprednisolone is a synthetic corticosteroid with anti-inflammatory and immunomodulating properties.



Purity: > 98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Methylprednisolone-d7

(U 7532-d7) Cat. No.: HY-B0260S3

Methylprednisolone-d7 is deuterium labeled Methylprednisolone. Methylprednisolone is a synthetic corticosteroid with anti-inflammatory and immunomodulating properties.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Metofenazate

(Methophenazine) Cat. No.: HY-100263

Metofenazate is a selective **calmodulin** inhibitor.



Purity: >98%

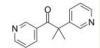
Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Metyrapone

(Su-4885) Cat. No.: HY-B1232

Metyrapone is an inhibitor of cytochrome P450-mediated ω/ω -1 hydroxylase activity and CYP11B1. Target: CYP11B1 Metyrapone is a drug used in the diagnosis of adrenal insufficiency and occasionally in the treatment of Cushing's syndrome (hypercortisolism).



Purity: 99.84% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg

Mevastatin

(Compactin; ML236B) Cat. No.: HY-17408

Mevastatin (Compactin) is a first HMG-CoA reductase inhibitor that belongs to the statins class. Mevastatin is a lipid-lowering agent, and induces apoptosis, arrests cancer cells in ${\rm G_0/G_1}$ phase.



Purity: 99.20%

Clinical Data: No Development Reported

Size: $10 \text{ mM} \times 1 \text{ mL}$, 50 mg, 100 mg, 500 mg

MG-132

(Z-Leu-Leu-al; MG132) Cat. No.: HY-13259

MG-132 (Z-Leu-Leu-Leu-al) is a potent proteasome and calpain inhibitor with IC $_{\rm S0}$ s of 100 nM and 1.2 μ M, respectively. MG-132 effectively blocks the proteolytic activity of the 26S proteasome complex. MG-132, a peptide aldehyde, also is an autophagy activator.



Purity: ≥98.0%

Clinical Data: No Development Reported

Size: 10 mg, 25 mg, 50 mg, 100 mg, 200 mg, 500 mg

MHY1485

Cat. No.: HY-B0795

MHY1485 is a potent cell-permeable mTOR activator that targets the ATP domain of mTOR. MHY1485 inhibits autophagy by suppression of fusion between autophagosomes and lysosomes.



Purity: 99.86%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

Mifepristone

(RU486; RU 38486)

Mifepristone (RU486) is a progesterone receptor (PR) and glucocorticoid receptor (GR) antagonist with IC_{s_0} s of 0.2 nM and 2.6 nM in in vitro assay.



Cat. No.: HY-13683

Purity: 99.77% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg

Mifepristone-13C,d3

(RU486-13C,d3; RU 38486-13C,d3)

Mifepristone-13C,d3 is the 13C- and deuterium labeled. Mifepristone (RU486) is a progesterone receptor (PR) and glucocorticoid receptor (GR)

antagonist with IC50s of 0.2 nM and 2.6 nM in in vitro assay.

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg



Cat. No.: HY-13683S1

Milciclib

(PHA-848125) Cat. No.: HY-10424

Milciclib (PHA-848125) is a potent, ATP-competitive and dual inhibitor of CDK and Tropomyosin receptor kinase (TRK), with IC_{so}s of 45, 150, 160, 363, 398 nM and 53 nM for cyclin A/CDK2, cyclin H/CDK7, cyclin D1/CDK4, cyclin E/CDK2, cyclin B/CDK1 and TRKA, respectively.

Purity: Clinical Data: Phase 2

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg Size:

Clinical Data: No Development Reported Size:

Purity:

Mirdametinib (PD0325901; PD325901)

Mifepristone-d3

(RU486-d3; RU 38486-d3)

and 2.6 nM in in vitro assay.

>98%

1 mg, 5 mg

Mifepristone-d3 (RU486-d3) is the deuterium

labeled Mifepristone, Mifepristone (RU486) is a

progesterone receptor (PR) and glucocorticoid receptor (GR) antagonist with IC₅₀s of 0.2 nM

Mirdametinib (PD0325901) is an orally active, selective and non-ATP-competitive MEK inhibitor with an IC_{so} of 0.33 nM. Mirdametinib exhibits a $\mathbf{K}_{i}^{\text{app}}$ of 1 nM against activated MEK1 and MEK2. Mirdametinib suppresses the expression of p-ERK1/2 and induces apoptosis.

MK-2206 is an orally active, highly potent and

respectively. Many breast cancer cell lines, and

PIK3CA-mutant and cell lines with PTEN loss are sensitive to MK-2206. Anticancer activities.

selective allosteric Akt inhibitor, with ICsos of

8, 12, and 65 nM for Akt1, Akt2, and Akt3,

Clinical Data: No Development Reported

1 mg, 5 mg

>98%

99 95% **Purity:** Clinical Data: Phase 2

MK-2206

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg



Mito-I ND

(Mito-Lonidamine) Cat. No.: HY-134832

Mito-LND (Mito-Lonidamine) is an orally active and mitochondria-targeted inhibitor of oxidative phosphorylation (OXPHOS). Mito-LND inhibits mitochondrial bioenergetics, stimulates the formation of reactive oxygen species, and induces autophagic cell death in lung cancer cells.

Purity: 97.00%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

MK-2206 dihydrochloride

(MK-2206 (2HCl)) Cat. No.: HY-10358

MK-2206 dihydrochloride (MK-2206 (2HCl)) is an orally active allosteric AKT inhibitor with IC₅₀s of 5 nM, 12 nM, and 65 nM for AKT1, AKT2, and AKT3, respectively. MK-2206 dihydrochloride induces autophagy.

Purity: 99.76% Clinical Data: Phase 2

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

MK-5108

Purity:

Size

(VX-689) Cat. No.: HY-13252

MK-5108 is a highly potent and specific inhibitor of Aurora A kinase with an IC_{so} value of 0.064 nM.

Cat. No.: HY-13683S

Cat. No.: HY-10254

Cat. No.: HY-108232

99.89% Purity: Clinical Data: Phase 1

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

ML327

Cat. No.: HY-103038

ML327 is a blocker of MYC which can also de-repress E-cadherin transcription and reverse Epithelial-to-Mesenchymal Transition (EMT).

Purity: 98.19%

No Development Reported Clinical Data:

Size 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

MLCK inhibitor peptide 18

Cat. No.: HY-P1029

MLCK inhibitor peptide 18 is a myosin light chain kinase (MLCK) inhibitor with an IC_{so} of 50 nM, and inhibits CaM kinase II only at 4000-fold higher concentrations.

RKKYKYRRK-NH2

Purity: 99.66%

Clinical Data: No Development Reported 1 mg, 5 mg, 10 mg, 25 mg

Mocetinostat

(MGCD0103) Cat. No.: HY-12164

Mocetinostat (MGCD0103) is a potent, orally active and isotype-selective HDAC (Class I/IV) inhibitor with IC₅₀s 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.



Purity: 99 43% Clinical Data: Phase 2

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg

Momelotinib

(CYT387) Cat. No.: HY-10961

Momelotinib (CYT387) is an ATP-competitive inhibitor of JAK1/JAK2 with IC and of 11 nM and 18 nM,respectively. CYT387 shows much less activity



98 93% Purity: Clinical Data: Phase 3

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

Momelotinib Mesylate

(CYT387 Mesylate) Cat. No.: HY-10963

Momelotinib Mesylate (CYT387 Mesylate) is an ATP-competitive inhibitor of JAK1/JAK2 with IC50 of 11 nM/18 nM, appr 10-fold selectivity versus JAK3.

Purity: >98% Clinical Data: Phase 3 1 mg, 5 mg

Momelotinib sulfate

(CYT387 sulfate salt) Cat. No.: HY-10962

Momelotinib sulfate (CYT387 sulfate salt) is an ATP-competitive inhibitor of JAK1/JAK2 with IC50 of 11 nM/18 nM, 10-fold selectivity versus JAK3 (IC₅₀=155 nM).

Purity: 98 04% Clinical Data: Phase 3

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Monacolin J

(Antibiotic MB 530A; Lovastatin diol lactone) Cat. No.: HY-104051

Monacolin J is an inhibitor of cholesterol biosynthesis, and inhibits the activity of HMG-CoA reductase.



>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

MRT67307

MRT67307 is a dual inhibitor of the IKKε and TBK-1 with IC₅₀s of 160 and 19 nM, respectively.

MRT67307 also inhibits ULK1 and ULK2 with IC_{so}s of 45 and 38 nM, respectively. MRT67307 also blocks autophagy in cells.

99.34% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

MRT67307 hydrochloride

Cat. No.: HY-13018A

MRT67307 hydrochloride is a dual inhibitor of the $IKK\epsilon$ and TBK-1 with $IC_{50}s$ of 160 and 19 nM, respectively. MRT67307 hydrochloride also inhibits ULK1 and ULK2 with IC_{so}s of 45 and 38 nM, respectively. MRT67307 hydrochloride also blocks autophagy in cells.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 ma, 5 ma

mTOR inhibitor-1

Cat. No.: HY-112914

mTOR inhibitor-1 is a novel mTOR pathway inhibitor which can suppress cells proliferation

and inducing autophagy.

Cat. No.: HY-13018

99.50% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

mTOR inhibitor-8

Cat. No.: HY-131344

mTOR inhibitor-8 is an mTOR inhibitor and autophagy inducer. mTOR inhibitor-8 inhibits the activity of mTOR via FKBP12 and induces autophagy of A549 human lung cancer cells.



Purity: 98.04%

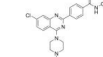
Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

mTOR/HDAC6-IN-1

Cat. No.: HY-144449

mTOR/HDAC6-IN-1 is a potent mTOR and HDAC6 dual inhibitor (IC $_{so}$ 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).



>98%

Clinical Data: No Development Reported

1 mg, 5 mg

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

MW-150

(MW01-18-150SRM) Cat. No.: HY-120111

MW150 (MW01-18-150SRM) is a selective, CNS penetrant, and orally active inhibitor of p38α MAPK with a K, of 101 nM. MW-150 inhibits the ability of the endogenous p38α MAPK to phosphorylate an endogenous substrate MK2 in activated glia.

Purity: 99 90%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

MW-150 hydrochloride

(MW01-18-150SRM hydrochloride) Cat. No.: HY-120111A

MW-150 hydrochloride (MW01-18-150SRM hydrochloride) is a selective, CNS penetrant, and orally active inhibitor of $p38\alpha$ MAPK with a K, of 101 nM.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Mycro 3

Purity:

Size:

Mycro 3 is a potent and selective inhibitor of Myc-associated factor X (MAX) dimerization. Mycro 3 also inhibit DNA binding of c-Myc. Mycro 3 could be used for the research of pancreatic cancer.

MW-150 dihydrochloride dihydrate (MW01-18-150SRM dihydrochloride dihydrate)

dihydrochloride dihydrate) is a selective. CNS

penetrant, and orally active inhibitor of $p38\alpha$

MAPK with a K, of 101 nM.

>98%

Clinical Data: No Development Reported

1 mg, 5 mg

MW-150 dihydrochloride dihydrate (MW01-18-150SRM

Purity: 99 21%

Clinical Data: No Development Reported

10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Myricetin

(Cannabiscetin) Cat. No.: HY-15097

Myricetin is a common plant-derived flavonoid with a wide range of activities including strong anti-oxidant, anticancer, antidiabetic and anti-inflammatory activities.

98.08% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 50 mg, 100 mg, 200 mg, 500 mg

N-Benzyloleamide

N-Benzyloleamide is a maccamide isolated from Lepidium meyenii (Maca). N-Benzyloleamide irreversibly inhibits fatty acid amide hydrolase (FAAH). N-benzyloleamide influences the energy metabolism and reveals antioxidant and antifatique

Cat. No.: HY-N6923

Cat. No.: HY-120111B

Cat. No.: HY-100669

Purity: 98.29%

Clinical Data: No Development Reported

Size: 5 mg, 10 mg

N-Benzylpalmitamide

(N-Benzylhexadecanamide; Macamide 1) Cat. No.: HY-N2365

N-Benzylpalmitamide is a macamide isolated from Lepidium meyenii, acts as an inhibitor of fatty acid amide hydrolase (FAAH).

98.39% Purity:

Clinical Data: No Development Reported

Size:

N-Benzyllinolenamide

N-Benzyllinolenamide is a natural macamide isolated from Lepidium meyenii, acts as an inhibitor of fatty acid amide hydrolase (FAAH)

with an IC_{50} of 41.8 μ M.

Cat. No.: HY-N3033

>98% Purity:

Clinical Data: No Development Reported 5 mg, 10 mg, 20 mg

N6-Isopentenyladenosine

(Riboprine) Cat. No.: HY-W011209

N6-Isopentenyladenosine (Riboprine), an RNA modification found in cytokinins, which regulate plant growth/differentiation, and a subset of tRNAs, where it improves the efficiency and accuracy of translation.



Purity: ≥98.0%

Clinical Data: No Development Reported

Size 25 mg, 50 mg

Nampt-IN-3

Cat. No.: HY-108701

Nampt-IN-3 (Compound 35) simultaneously inhibit nicotinamide phosphoribosyltransferase (NAMPT) and HDAC with ICsos of 31 nM and 55 nM, respectively. Nampt-IN-3 effectively induces cell apoptosis and autophagy and ultimately leads to cell death.



Purity: 99.27%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Naproxen

((S)-Naproxen) Cat. No.: HY-15030

Naproxen is a COX-1 and COX-2 inhibitor with IC_{so} s of 8.72 and 5.15 μ M, respectively in cell assay.

99 98% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 5 g, 10 g

Naproxen sodium

Naproxen sodium is a COX-1 and COX-2 inhibitor with IC_{so}s of 8.72 and 5.15 μM, respectively in cell assay.



Cat. No.: HY-15030A

99 98% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 5 g, 10 g

Naringin

(Naringoside) Cat. No.: HY-N0153

Naringin is a major flavanone glycoside obtained from tomatoes, grapefruits, and many other citrus fruits. Naringin exhibits biological properties such as antioxidant, anti-inflammatory, and antiapoptotic activities.

Purity: 98.44%

Clinical Data: No Development Reported 10 mM × 1 mL, 200 mg, 10 g

NBDHEX

Cat. No.: HY-135318

NBDHEX is a potent glutathione S-transferase P1-1 (GSTP1-1) inhibitor. NBDHEX induces apoptosis of tumor cells.



Purity: 98.58%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg Size:

Necrostatin-1

(Nec-1) Cat. No.: HY-15760

Necrostatin-1 (Nec-1) is a potent necroptosis inhibitor with an EC₅₀ of 490 nM in Jurkat cells. Necrostatin-1 inhibits RIP1 kinase (EC_{so}=182 nM). Necrostatin-1 is also an IDO inhibitor.



Purity: 99.87%

Clinical Data: No Development Reported

10 mM × 1 mL, 10 mg, 50 mg, 100 mg Size:

Neferine

((-)-Neferine) Cat. No.: HY-N0441

Neferine is a major bisbenzylisoquinline alkaloid. Neferine strongly inhibits NF-κB activation.



99.92% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg Size:

Neflamapimod

(VX-745) Cat. No.: HY-10328

Neflamapimod (VX-745) is a potent, blood-brain barrier penetrant, highly selective inhibitor of $p38\alpha$ inhibitor with an IC_{so} for $p38\alpha$ of 10 nM and for p38β of 220 nM. Neflamapimod (VX-745) possesses anti-inflammatory activity.



99.32% Purity: Clinical Data: Phase 2

10 mM × 1 mL, 10 mg, 50 mg Size:

Nesolicaftor

(PTI-428) Cat. No.: HY-111680

Nesolicaftor (PTI-428) is a specific cystic fibrosis transmembrane conductance regulator (CFTR) amplifier.

Purity: 99.65% Clinical Data: Phase 2

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

NH125

Cat. No.: HY-100576

NH125 is a potent and selective inhibitor of eukaryotic elongation factor 2 kinase (eEF-2K/CaMKIII), also can induce eEF2 phosphorylation, with an IC₅₀ of 60 nM for eEF-2K.



Purity: ≥98.0%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Niacin

(Nicotinic acid; Vitamin B3)

Niacin (Nicotinic acid) is a vitamin and is part

of the vitamin B group.



Cat. No.: HY-B0143

99.77% Purity: Clinical Data: Launched

10 mM × 1 mL, 500 mg, 1 g, 5 g Size:

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

Niacin-13C6

(Nicotinic acid-13C6; Vitamin B3-13C6)

Niacin-13C6 (Nicotinic acid-13C6) is the 13C-labeled Niacin. Niacin (Nicotinic acid) is a vitamin and is part of the vitamin B group.

Purity: > 98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Cat. No.: HY-B0143S3 Nicardipine hydrochloride (YC-93)

(YC-93) Cat. No.: HY-12515A

Nicardipine hydrochloride (YC-93) is a **calcium channel** blocker with an IC_{50} of 1 μ M for blocking cardiac calcium channels. Nicardipine hydrochloride acts as an agent for chronic stable angina and for controlling blood pressure.

HN H-GI

Cat. No.: HY-17640

Purity: 99.72% Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g

Niclosamide olamine

(BAY2353 olamine) Cat. No.: HY-B0497C

Niclosamide olamine (BAY2353 olamine) is an anthelmintic that disrupts mitochondrial metabolism in parasitic worms and animal models.

Purity: >98%
Clinical Data: Phase 4
Size: 1 mg, 5 mg

Nicodicosapent

Nicodicosapent is a fatty acid niacin conjugate that is also an inhibitor of the sterol regulatory element binding protein (SREBP), a key regulator of cholesterol metabolism proteins such as PCSK9,

HMG-CoA reductase, ATP citrate lyase, and NPC1L1.

Purity: 98.04%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 50 mg, 100 mg

Nidufexor

(LMB763) Cat. No.: HY-109096

Nidufexor (LMB763) is an orally-available **farnesoid** X **receptor** (FXR) agonist for the research of nonalcoholic steatohepatitis (NASH).

Purity: 98.96%

Clinical Data: No Development Reported

Size: $10 \text{ mM} \times 1 \text{ mL}$, 5 mg, 10 mg, 50 mg, 100 mg

Nifedipine

(BAY-a-1040) Cat. No.: HY-B0284

Nifedipine (BAY-a-1040) is a potent **calcium channel** blocker and drug of choice for cardiac insufficiencies.



Purity: 99.35% Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g, 10 g

Nifedipine-d4

(BAY-a-1040-d4) Cat. No.: HY-B0284S1

Nifedipine-d4 (BAY-a-1040-d4) is the deuterium labeled Nifedipine. Nifedipine (BAY-a-1040) is a potent **calcium channel** blocker and drug of choice for cardiac insufficiencies.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Nilotinib (AMN107)

Nilotinib is an orally available **Bcr-Abl** tyrosine kinase inhibitor with antineoplastic activity.

Cat. No.: HY-10159

Purity: 99.96% Clinical Data: Launched

Size: 100 mg, 200 mg, 500 mg

Nilotinib monohydrochloride monohydrate

(AMN107 monohydrochloride monohydrate) Cat. No.: HY-10159A

Nilotinib monohydrochloride monohydrate is a second generation tyrosine kinase inhibitor (TKI), is significantly potent against BCR-ABL, and is active against many BCR-ABL mutants.

Purity: 99.89% Clinical Data: Launched

Size: $10 \text{ mM} \times 1 \text{ mL}$, 100 mg, 200 mg, 500 mg

Nilotinib-d3

Cat. No.: HY-132549S

Nilotinib-d3 (AMN107-d3) is the deuterium labeled Nilotinib. Nilotinib is an orally available **Bcr-Abl** tyrosine kinase inhibitor with antineoplastic activity.



Purity: >98%

Clinical Data:

Size: 1 mg, 10 mg

Nilotinib-d6

(AMN107-d6) Cat. No.: HY-10159S

Nilotinib D6 (AMN107 D6) is a deuterium labeled Nilotinib. Nilotinib is an orally available Bcr-Abl tyrosine kinase inhibitor with antineoplastic activity.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Nimodipine-d7

Cat. No.: HY-B0265S Nimodipine-d7 is the deuterium labeled Nimodipine.

Nimodipine (BAY-e 9736) is an orally active, well-tolerated and light-sensitive dihydropyridine calcium antagonist. Nimodipine can be used for the research of cerebrovascular disorders.

Purity: >98%

Clinical Data: No Development Reported 1 mg, 2 mg, 5 mg, 10 mg Size:

Nitrendipine (BAY-E-5009) Cat. No.: HY-B0424

Nitrendipine (BAY-E-5009), an analogue of Nifedipine (HY-B0284), is a dihydropyridine calcium channel blocker with vasodilator action. Nitrendipine has antihypertensive effect.

Purity: 99 25% Clinical Data: Launched

Size: 10 mM × 1 mL, 50 mg, 100 mg, 200 mg, 500 mg, 1 g

Nitroprusside disodium dihydrate (Sodium nitroprusside

dihydrate; Sodium Nitroferricyanide(III) Dihydrate) Cat. No.: HY-A0119

Nitroprusside disodium dehydrate (Sodium nitroprusside dihydrate) is a vasodilator that available for the research of acute hypertension, heart failure. Nitroprusside disodium dehydrate induces autophagy in glutathione-depleted osteoblasts.

Purity: 99.72% Clinical Data: Launched

10 mM × 1 mL, 500 mg, 10 g Size

Na₂[Fe(CN)₅NO].2H₂O

Nitroxoline-D4

(8-Hydroxy-5-nitroquinoline-D4; 5-Nitro-8-quinolinol-D4) Cat. No.: HY-B1159S

Nitroxoline-D4 (8-Hydroxy-5-nitroquinoline-D4) is the deuterium labeled Nitroxoline. Nitroxoline is an antibiotic that has proven to be very effective at combating biofilm infections. Nitroxoline functions by chelating Fe2+ and Zn2+ ions from the biofilm matrix.

Purity: >98%

Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg

Nimodipine

(BAY-e 9736) Cat. No.: HY-B0265

Nimodipine (BAY-e 9736) is an orally active, well-tolerated and light-sensitive dihydropyridine calcium antagonist. Nimodipine can be used for the research of cerebrovascular disorders.

Purity: 99 76% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg

Nitazoxanide

(NTZ; NSC 697855)

Nitazoxanide (NTZ), an anthelmintic agent, exhibits a broad spectrum of activities against a wide variety of helminths, protozoa, and enteric bacteria infecting animals and humans.

Cat. No.: HY-B0217

Purity: 98 35% Clinical Data: Launched

10 mM × 1 mL, 10 mg, 50 mg, 100 mg

Nitrendipine-d5

(AY-E-5009-d5) Cat. No.: HY-B0424S

Nitrendipine-d5 (AY-E-5009-d5) is the deuterium labeled Nitrendipine. Nitrendipine (BAY-E-5009), an analogue of Nifedipine (HY-B0284), is a dihydropyridine calcium channel blocker with vasodilator action. Nitrendipine has antihypertensive effect.

Purity: >98%

Clinical Data: No Development Reported

Size 1 mg, 5 mg

Nitroxoline

(8-Hydroxy-5-nitroquinoline; 5-Nitro-8-quinolinol) Cat. No.: HY-B1159

Nitroxoline is an antibiotic that has proven to be very effective at combating biofilm infections. Nitroxoline functions by chelating Fe2+ and Zn2+ ions from the biofilm matrix.

99.57% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 1 g

NL-1

Cat. No.: HY-135231

NL-1 is a mitoNEET inhibitor with antileukemic effect. NL-1 inhibits REH and REH/Ara-C cells growth with IC_{so}s of 47.35 µM and 56.26 µM, respectively. NL-1-mediated death in leukemic cells requires the activation of the autophagic pathway.

Purity: 99.11%

Clinical Data: No Development Reported 10 mM × 1 mL, 10 mg, 50 mg

Nobiletin

Cat. No.: HY-N0155

Nobiletin is a poly-methoxylated flavone from the citrus peel that improves memory loss. Nobiletin is a retinoid acid receptor-related orphan receptors (RORs) agonist.

99 52% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Norepinephrine

Purity:

Size:

Nocodazole

(Oncodazole; R17934)

(Levarterenol; L-Noradrenaline)

Norepinephrine (Levarterenol; L-Noradrenaline) is a potent adrenergic receptor (AR) agonist. Norepinephrine activates α_1 , α_2 , β_1 receptors.

Nocodazole (Oncodazole) is a rapidly-reversible inhibitor of microtubule. Nocodazole binds to

assembly/disassembly dynamics, which prevents mitosis and induces apoptosis in tumor cells.

β-tubulin and disrupts microtubule

99 66%

Clinical Data: No Development Reported

Cat. No.: HY-13715

Cat. No.: HY-13520

Purity: 98.08% Clinical Data: Launched 500 ma

Nordihydroguaiaretic acid

(NDGA) Cat. No.: HY-N0198

Nordihydroguaiaretic acid is a 5-lipoxygenase (5LOX) (IC₅₀=8 μ M) and tyrosine kinase inhibitor.

Purity: 99 88% Clinical Data: Phase 2

10 mM × 1 mL, 100 mg, 250 mg

Norepinephrine bitartrate monohydrate (Levarterenol

bitartrate monohydrate; ...) Cat. No.: HY-13715B

Norepinephrine (Levarterenol; L-Noradrenaline) bitartrate monohydrate is a potent adrenergic receptor (AR) agonist. Norepinephrine activates α_1 , α_2 , β_1 receptors.

99.75% Purity: Clinical Data: Launched Size: 500 mg, 1 g, 5 g

Norepinephrine hydrochloride (Levarterenol hydrochloride;

10 mM × 1 mL, 10 mg, 50 mg, 100 mg

L-Noradrenaline hydrochloride)

Norepinephrine (Levarterenol; L-Noradrenaline) hydrochloride is a potent adrenergic receptor (AR) agonist. Norepinephrine activates α_1 , α_2 , β_1 receptors.

HCI

Cat. No.: HY-13715A

Purity: 99.95% Clinical Data: Launched Size 500 ma

Norepinephrine tartrate

(Levarterenol tartrate; L-Noradrenaline tartrate) Cat. No.: HY-13715C

Norepinephrine (Levarterenol; L-Noradrenaline) tartrate is a potent adrenergic receptor (AR) agonist. Norepinephrine tartrate activates α_{ij} α_{2} , β_{1} receptors.

>98% Purity: Clinical Data: Launched Size 5 mg, 10 mg, 25 mg

Norswertianin

Norswertianin, a xanthone compound, serves as a powerful anti-glioma compound. Norswertianin induces GBM cells differentiation through oxidative stress and Akt/mTOR dependent autophagy.

Cat. No.: HY-N9341

>98% Purity:

Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg

Nortriptyline

(Desmethylamitriptyline; Desitriptilina) Cat. No.: HY-118620

Nortriptyline (Desmethylamitriptyline), the main active metabolite of Amitriptyline, is a tricyclic antidepressant used to relieve the symptoms of depression. Nortriptyline is a potent autophagy inhibitor.

Purity: >98% Clinical Data: Launched

Size 10 mg, 50 mg, 100 mg

Nortriptyline hydrochloride (Desmethylamitriptyline

hydrochloride; Desitriptilina hydrochloride)

Nortriptyline hydrochloride (Desmethylamitriptyline hydrochloride), the main active metabolite of Amitriptyline, is a tricyclic antidepressant used to relieve the symptoms of depression. Nortriptyline hydrochloride is a potent autophagy inhibitor.

Purity: 99.96% Clinical Data: Launched

10 mM × 1 mL, 50 mg

Cat. No.: HY-B1417

Nortriptyline-d3 hydrochloride (Desmethylamitriptyline-d3

hydrochloride; Desitriptilina-d3 hydrochloride) Cat. No.: HY-B1417S

Nortriptyline-d3 (Desmethylamitriptyline-d3) hydrochloride is the deuterium labeled Nortriptyline hydrochloride.

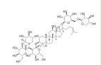


>98% Purity:

Clinical Data: No Development Reported Size: 2.5 mg, 1 mg, 5 mg, 10 mg

Notoginsenoside Fc

Notoginsenoside Fc, a protopanaxadiol- (PPD-) type saponin isolated from the leaves of Panax notoginseng, effectively counteracts platelet aggregation. Notoginsenoside Fc can accelerate reendothelialization following vascular injury in diabetic rats by promoting autophagy.



Cat. No.: HY-16916

Cat. No.: HY-N2531

Purity: >98%

Clinical Data: No Development Reported

Size: 5 mg, 10 mg

Novobiocin Sodium

(Albamycin; Cathomycin) Cat. No.: HY-B0425A

Novobiocin Sodium (Albamycin; Cathomycin) is an orally active antibiotic compound derived from Streptomyces niveus and a potent DNA gyrase inhibitor by binding the ATP-binding site in the ATPase subunit.



Purity: 99.12% Clinical Data: Launched

10 mM × 1 mL, 100 mg, 500 mg

NS1643

NS1643 is a partial agonist of human ether-a-go-go-related gene (hERG) K(+) channels with an EC_{so} of 10.5 μ M. NS1643 has distinct effects on erg2 (Kv11.2) currents by reducing channel inactivation especially at high concentrations.

Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 500 mg

NSC 185058

Cat. No.: HY-125169

NSC 185058 is an inhibitor of ATG4B, a major cysteine protease. Inhibition of ATG4B using NSC 185058 markedly attenuates autophagic activity.

99.52% Purity:

Clinical Data: No Development Reported

Size: 5 mg, 10 mg

NSC 33994

Cat. No.: HY-18293

NSC 33994 (G6) is a selective JAK2 inhibitor, with an IC₅₀ of 60 nM.



>98% Purity:

Clinical Data: No Development Reported

Size 1 mg, 5 mg

Nutlin-3a

(Rebemadlin) Cat. No.: HY-10029

Nutlin-3a (Rebemadlin), an active enantiomer of Nutlin-3, is a potent murine double minute (MDM2) inhibitor (IC_{so}=90 nM). Nutlin-3a inhibits MDM2-p53 interactions and stabilizes the p53 protein, and induces cell autophagy and apoptosis.



Purity: 98.07%

Clinical Data: No Development Reported 10 mM \times 1 mL, 5 mg, 10 mg, 50 mg Size

NVP-AEW541

(AEW541) Cat. No.: HY-50866

NVP-AEW541 (AEW541) is a potent inhibitor of IGF-1R with IC_{50} of 0.15 μM , also inhibits InsR, with IC_{50}

of 0.14 μM.

Purity: 98.90%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 50 mg, 100 mg

Obatoclax

(GX15-070) Cat. No.: HY-10969A

Obatoclax (GX15-070), a BH3 mimetic, is a pan-BCL-2 family proteins inhibitor with a K, of 220 nM for BCL-2. Obatoclax induces autophagy-dependent cell death and targets cyclin D1 for proteasomal degradation.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Obatoclax Mesylate

(GX15-070 Mesylate)

Obatoclax Mesylate (GX15-070 Mesylate), a BH3 mimetic, is a pan-BCL-2 family proteins inhibitor with a K, of 220 nM for BCL-2. Obatoclax Mesylate induces autophagy-dependent cell death and targets cyclin D1 for proteasomal degradation.



Cat. No.: HY-10969

99.74% Clinical Data: Phase 3

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

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Obeticholic acid

(INT-747; 6-ECDCA; 6-Ethylchenodeoxycholic acid) Cat. No.: HY-12222

Obeticholic acid (INT-747) is a potent, selective and orally active FXR agonist with an EC_{50} of 99 nM. Obeticholic acid has anticholeretic and anti-inflammation effect. Obeticholic acid also induces autophagy.



Purity: ≥98.0% Clinical Data: Launched

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

Obeticholic Acid-d4

Obeticholic Acid-d4 is the deuterium labeled Obeticholic acid. Obeticholic acid (INT-747) is a potent, selective and orally active FXR agonist with an EC_{50} of 99 nM. Obeticholic acid has anticholeretic and anti-inflammation effect. Obeticholic acid also induces **autophagy**.



Cat. No.: HY-12222S1

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Obeticholic acid-d5

(INT-747-d5; 6-ECDCA-d5; 6-Ethylchenodeoxycholic acid-d5) Cat. No.: HY-12222S

Obeticholic acid-d5 (INT-747-d5) is the deuterium labeled Obeticholic acid. Obeticholic acid (INT-747) is a potent, selective and orally active FXR agonist with an EC $_{50}$ of 99 nM. Obeticholic acid has anticholeretic and anti-inflammation effect. Obeticholic acid also induces autophagy.



Purity: >98%

Clinical Data: No Development Reported

Size: 2.5 mg, 1 mg

Olacaftor

(VX-440) Cat. No.: HY-112267

Olacaftor (VX-440) is a cystic fibrosis transmembrane conductance regulator (CFTR) modulator extracted from patent US9782408.



Purity: >98%

Clinical Data: No Development Reported

Size: 5 mg, 10 mg

Olaparib

(AZD2281; KU0059436) Cat. No.: HY-10162

Olaparib (AZD2281; KU0059436) is a potent and orally active PARP inhibitor with ${\rm IC}_{\rm so}{\rm S}$ of 5 and 1 nM for PARP1 and PARP2, respectively. Olaparib is an autophagy and mitophagy activator.



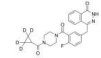
Purity: 99.98% Clinical Data: Launched

Size: $10 \text{ mM} \times 1 \text{ mL}$, 10 mg, 50 mg, 100 mg, 200 mg, 500 mg

Olaparib-d4-1

(AZD2281-d4-1; KU0059436-d4-1) Cat. No.: HY-10162S3

Olaparib-d4-1 (AZD2281-d4-1) is the deuterium labeled Olaparib. Olaparib (AZD2281; KU0059436) is a potent and orally active PARP inhibitor with $\rm IC_{so}S$ of 5 and 1 nM for PARP1 and PARP2, respectively. Olaparib is an autophagy and mitophagy activator.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Olaparib-d5

(AZD2281-d5; KU0059436-d5) Cat. No.: HY-10162S

Olaparib D5 (AZD2281 D5) is a deuterium labeled Olaparib. Olaparib is a potent and oral PARP inhibitor.



Purity: > 98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Olaparib-d8

(AZD2281-d8; KU0059436-d8) Cat. No.: HY-10162S1

Olaparib D8 (AZD2281 D8) is the deuterium labeled Olaparib (AZD2281). Olaparib is a potent and orally active PARP inhibitor with $\rm IC_{50}$ s of 5 and 1 nM for PARP1 and PARP2, respectively. Olaparib is an <code>autophagy</code> and <code>mitophagy</code> activator.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Oleanolic Acid

(Oleanic acid; Caryophyllin) Cat. No.: HY-N0156

Oleanolic acid (Caryophyllin) is a natural compound from plants with anti-tumor activities.



Purity: ≥98.0%
Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg

Omeprazole

(H 16868) Cat. No.: HY-B0113

Omeprazole (H 16868), a **proton pump** inhibitor (PPI), is available for treatment of acid-related gastrointestinal disorders. Omeprazole shows competitive inhibition of **CYP2C19** activity with a $K_{\rm i}$ of 2 to 6 μM .



Purity: 98.19% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg

Omeprazole sodium

(H 16868 sodium) Cat. No.: HY-B0113A

Omeprazole sodium (H 16868 sodium), a **proton pump** inhibitor (PPI), is available for treatment of acid-related gastrointestinal disorders. Omeprazole sodium shows competitive inhibition of **CYP2C19** activity with a \mathbf{K}_i of 2 to 6 μ M.

Na Na

Purity: 98.03% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg

Omeprazole-13CD3

(H 16868-13CD3) Cat. No.: HY-B0113S3

Omeprazole-13CD3 (H 16868-13CD3) is a 13C-labeled and deuterium labeled Omeprazole. Omeprazole (H 16868), a **proton pump** inhibitor (PPI), is available for treatment of acid-related gastrointestinal disorders.

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Purity: >98%

Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg

512c. 1 mg, 5 mg, 10 mg

Omeprazole-d3

(H 16868-d3) Cat. No.: HY-B0113S

Omeprazole D3 (H 16868 D3) is deuterium labeled Omeprazole. Omeprazole, a **proton pump** inhibitor (PPI), is available for treatment of acid-related gastrointestinal disorders.

Purity: 98.99%

Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg

Omeprazole-d3-1

(H 16868-d3-1) Cat. No.: HY-B0113S1

Omeprazole-d3-1 (H 16868-d3-1) is the deuterium labeled Omeprazole. Omeprazole (H 16868), a proton pump inhibitor (PPI), is available for treatment of acid-related gastrointestinal disorders.



Purity: >98%

Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg

Omipalisib

(GSK2126458; GSK458) Cat. No.: HY-10297

Omipalisib (GSK2126458) is an orally active and highly selective inhibitor of PI3K with K₁s of 0.019 nM/0.13 nM/0.024 nM/0.06 nM and 0.18 nM/0.3 nM for p110c/ β / δ / γ , mTORC1/2, respectively. Omipalisib has anti-cancer activity.



99 93%

Purity: 99.93% Clinical Data: Phase 1

Size: $10 \text{ mM} \times 1 \text{ mL}$, 5 mg, 10 mg, 50 mg

Onjisaponin B

Onjisaponin B is a natural product derived from Radix Polygalae. Onjisaponin B enhances autophagy and accelerates the degradation of mutant $\alpha\text{-synuclein}$ and huntingtin in PC-12 cells, and exbibits potential therapeutic effects on Parkinson disease and Huntington disease.



Cat. No.: HY-N2099

Purity: 99.10%

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 5 mg, 10 mg

Oprozomib

(ONX 0912; PR-047) Cat. No.: HY-12113

Oprozomib (PR-047) is an orally bioavailable and selective peptide epoxyketone **proteasome** inhibitor with \mathbf{IC}_{sg} s of 36 and 82 nM for proteasome (β 5) and immunoproteasome (LMP7), respectively. Oprozomib (ONX 0912) induces apoptosis in MM cells.



Purity: 99.71% Clinical Data: Phase 2

Size: $10 \text{ mM} \times 1 \text{ mL}$, 5 mg, 10 mg, 50 mg, 100 mg

OR-1855

OR-1855, an active metabolite of Levosimendan, has effect on human myometrial contractility. Levosimendan is a calcium sensitiser used in the management of acutely decompensated congestive heart failure



Cat. No.: HY-W050000

Purity: ≥97.0%

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 100 mg

Oroxin B

Cat. No.: HY-N1435

Oroxin B (OB) is a flavonoid isolated from traditional Chinese herbal medicine Oroxylum indicum (L.) Vent.



Purity: 99.71%

Clinical Data: No Development Reported

Size: 5 mg, 10 mg

Oroxylin A

(Baicalein 6-methyl ether; 6-Methoxybaicalein)

Oroxylin A is a natural active flavonoid with strong anticancer effects. IC50 value: Target: In vitro: Oroxylin A suppressed the MDM2-mediated degradation of p53 via downregulating MDM2 transcription in wt-p53 cancer cells .



Cat. No.: HY-N0560

Purity: 99.90%

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 5 mg, 10 mg

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OSI-027

(ASP7486) Cat. No.: HY-10423

OSI-027 (ASP7486) is a potent, selective, orally active and ATP-competitive mTOR kinase activity inhibitor with an IC_{50} of 4 nM. OSI-027 targets both mTORC1 and mTORC2 with IC_{so}s of 22 nM and 65 nM, respectively.

Purity: 99 40% Clinical Data: Phase 1

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg OSU-T315

OSU-T315 (ILK-IN-1) is a small Integrin-linked kinase (ILK) inhibitor with an IC₅₀ of 0.6 μM, inhibiting PI3K/AKT signaling by dephosphorylation of AKT-Ser473 and other ILK targets (GSK-3β and myosin light chain).

Cat. No.: HY-18676

99.88% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg

Ouabain Octahydrate

(Acocantherine; G-Strophanthin) Cat. No.: HY-B0542

Ouabain Octahydrate is an inhibitor of Na+/K+-ATPase, used for the treatment of congestive heart failure.

Purity: 99 96% Clinical Data: Launched

10 mM × 1 mL, 100 mg Size:

Oxaliplatin

Cat. No.: HY-17371

Oxaliplatin is a DNA synthesis inhibitor. Oxaliplatin causes DNA crosslinking damage, prevents DNA replication and transcription and causes cell death.

Purity: 99.57% Clinical Data: Launched

Oxidopamine hydrochloride

an antagonist of the neurotransmitter

dopamine, is a widely used neurotoxin that

selectively destroys dopaminergic neurons.

5 mg, 50 mg, 100 mg, 200 mg, 500 mg

Oxidopamine hydrochloride (6-OHDA hydrochloride),

(6-Hydroxydopamine hydrochloride; 6-OHDA hydrochloride) Cat. No.: HY-B1081

Oxidopamine hydrobromide

(6-Hydroxydopamine hydrobromide; 6-OHDA hydrobromide)Cat. No.: HY-B1081A

Oxidopamine hydrobromide (6-OHDA hydrobromide),

an antagonist of the neurotransmitter dopamine, is a widely used neurotoxin that selectively destroys dopaminergic neurons.

H-Br

Purity: 99.65%

Clinical Data: No Development Reported Size: 50 mg, 100 mg, 200 mg, 500 mg, 1 g

≥98.0% Purity:

Clinical Data: No Development Reported

Size 5 mg

Oxidopamine-d4 hydrobromide

(6-Hydroxydopamine-d4 hydrobromide; 6-OHDAd4 hydrobromide): HY-B1081AS

Oxidopamine-d4 (6-Hydroxydopamine-d4) hydrobromide is the deuterium labeled Oxidopamine hydrobromide. Oxidopamine (6-OHDA) hydrobromide,

an antagonist of the neurotransmitter dopamine, is a widely used neurotoxin that selectively destroys dopaminergic neurons.

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg Oxybenzone (Benzophenone 3)

Oxybenzone (Benzophenone 3) is a commonly used UV

filter in sun tans and skin protectants.

Oxybenzone act as endocrine disrupting chemicals (EDCs) and can pass through the placental and

blood-brain barriers.

99.91% Purity: Clinical Data: Launched

10 mM × 1 mL, 500 mg, 5 g Size:

Cat. No.: HY-A0067

H-CI

Oxyphenisatin acetate

Cat. No.: HY-101714

Oxyphenisatin acetate, the pro-drug of oxyphenisatin, is used to be a laxative.

Purity: ≥98.0%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg Oxyresveratrol

(trans-Oxyresveratrol)

Oxyresveratrol (trans-Oxyresveratrol) is a potent naturally occurring antioxidant and free radical scavenger (IC₅₀ of 28.9 μM against DPPH free

radicals).

98.87%

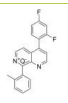
Clinical Data: No Development Reported 10 mM × 1 mL, 100 mg, 500 mg, 1 g

Cat. No.: HY-N1430

p38 MAPK-IN-1

Cat. No.: HY-12839

p38 MAPK-IN-1 (Compound 4) is a novel potent and selective inhibitor of p38 MAPK with IC $_{50}$ of 68 nM. p38 MAPK-IN-1 shows sustained levels, low clearance and good bioavailability.



Purity: 98.91%

Clinical Data: No Development Reported
Size: 5 mg, 10 mg, 50 mg, 100 mg

p38 MAPK-IN-2

p38 MAPK-IN-2 is an inhibitor of p38 kinase.



Cat. No.: HY-U00324

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

p38-α MAPK-IN-1

Cat. No.: HY-18874

p38- α MAPK-IN-1 is an inhibitor of MAPK14 (p38- α), with IC $_{s0}$ of 2300 nM in EFC displacement assay, and 5500 nM in HTRF assay.



Purity: 99.90%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 50 mg, 100 mg

p38α inhibitor 1

Cat. No.: HY-114423

p38 α inhibitor 1 is a p38 α inhibitor extracted from patent WO 2008076265 A1.



Purity: 98.70%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

PAC-1

(Procaspase activating compound 1)

apoptosis in cancer cells with an EC₅₀ of 2.08

PAC-1 is a **procaspase-3** activator that induces

μΜ.

orang.

Cat. No.: HY-13523

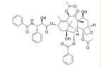
Purity: 99.93% Clinical Data: Phase 2

Size: 10 mM \times 1 mL, 10 mg, 50 mg, 100 mg, 500 mg

Paclitaxel

Cat. No.: HY-B0015

Paclitaxel is a naturally occurring antineoplastic agent and stabilizes **tubulin** polymerization. Paclitaxel can cause both mitotic arrest and **apoptotic** cell death. Paclitaxel also induces **autophagy**.



Purity: 99.97% Clinical Data: Launched

Size: 10 mM × 1 mL, 50 mg, 100 mg, 500 mg

Paclitaxel-d5 (benzoyloxy)

Cat. No.: HY-B0015S1

Paclitaxel-d5 benzoyloxy is the deuterium labeled Paclitaxel. Paclitaxel is a naturally occurring antineoplastic agent and stabilizes tubulin polymerization. Paclitaxel can cause both mitotic arrest and apoptotic cell death. Paclitaxel also induces autophagy.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Paeonol

Cat. No.: HY-N0159

Paeonol is an active extraction from the root of Paeonia suffruticosa, Paeonol inhibits MAO-A and MAO-B with IC $_{50}$ of 54.6 μM and 42.5 $\mu\text{M},$ respectively.

Purity: 99.86% Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 1 g



Palovarotene

(R 667; Ro 3300074) Cat. No.: HY-14799

Palovarotene is a nuclear retinoic acid receptor γ (RAR- $\!\gamma\!$) agonist.



Purity: 99.49% Clinical Data: Phase 3

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Pamapimod

(Ro4402257; R1503)

Pamapimod (Ro4402257) is a potent, selective and orally active **p38 MAPK** inhibitor with IC_{so} s of 14 nM and 480 nM and K_i s of 1.3 nM and 120 nM for **p38\alpha** and **p38\beta**, respectively. Pamapimod has no activity against p38 δ or p38 γ isoforms.



Cat. No.: HY-10405

Purity: 99.92% Clinical Data: Launched

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg

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

Pamapimod-d4

Pamapimod-d4 (Ro4402257-d4) is the deuterium labeled Pamapimod, Pamapimod (Ro4402257) is a potent, selective and orally active p38 MAPK inhibitor with IC₅₀s of 14 nM and 480 nM and K_is of 1.3 nM and 120 nM for p38 α and p38 β , respectively.

Purity: >98%

Clinical Data:

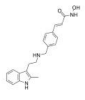
Size: 1 mg, 5 mg, 10 mg

Cat. No.: HY-10405S

(LBH589; NVP-LBH589)

Panobinostat

Panobinostat (LBH589; NVP-LBH589) is a potent and orally active non-selective HDAC inhibitor, and has antineoplastic activities.



Cat. No.: HY-10224

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)

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.



Cat. No.: HY-10224S

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Panobinostat-d4 hydrochloride

(LBH589-d4 hydrochloride; NVP-LBH589-d4 hydrochloride) Cat. No.: HY-10224S1

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

1 mg, 5 mg

Pantoprazole

(BY1023; SKF96022) Cat. No.: HY-17507

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 ICso of 6.8 μM.

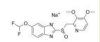


Purity: >98% Clinical Data: Launched Size: 1 mg, 5 mg

Pantoprazole sodium

(BY1023 sodium; SKF96022 sodium)

Pantoprazole sodium (BY10232 sodium) is an orally active and potent proton pump inhibitor (PPI). Pantoprazole sodium, a substituted benzimidazole, is a potent H+/K+-ATPase inhibitor with an IC_{so}



Cat. No.: HY-17507A

Purity: 99 89% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg

Pantoprazole sodium hydrate

(BY1023 sodium hydrate; SKF96022 sodium hydrate) Cat. No.: HY-17507B

Pantoprazole sodium hydrate (BY10232 sodium hydrate) is an orally active and potent proton pump inhibitor (PPI). Pantoprazole sodium hydrate, a substituted benzimidazole, is a potent H^+/K^+ -ATPase inhibitor with an IC_{50} of 6.8 μM .



99.94% Purity: Clinical Data: Launched

10 mM × 1 mL, 100 mg, 500 mg Size:

Pantoprazole-d3

(BY1023-d3; SKF96022-d3)

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+-ATP as inhibitor with an IC50 of 6.8 μ M.



Cat. No.: HY-17507S1

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Pantoprazole-d6

(BY1023-d6; SKF96022-d6) Cat. No.: HY-17507S

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 μ M.



Purity: >98%

Clinical Data: No Development Reported

Size 1 mg, 5 mg

Paris saponin VII

(Chonglou Saponin VII)

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-qp.



Cat. No.: HY-N3584

Purity: 99.13%

Clinical Data: No Development Reported

5 mg, 10 mg

Paroxetine hydrochloride

(BRL29060 hydrochloride; BRL29060A)

Paroxetine hydrochloride is a potent selective serotonin-reuptake inhibitor, commonly prescribed as an and has GRK2 inhibitory ability with IC50 of 14μM. Paroxetine hydrochloride can be used for the research of depressive disorder.

Cat. No.: HY-B0492

Purity: 99 92% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg

Paroxetine hydrochloride hemihydrate (BRL29060 hydrochloride

hemihydrate; BRL29060A hemihydrate)

Paroxetine hydrochloride hemihydrate is a potent selective serotonin-reuptake inhibitor, commonly prescribed as an antidepressant and has GRK2 inhibitory ability with IC₅₀ of 14μM.



Cat. No.: HY-B0492A

>98% Purity: Clinical Data: Launched Size: 1 mg, 5 mg

Paroxetine-d4 hydrochloride

(BRL29060-d4 hydrochloride; BRL29060A-d4)

Paroxetine-d4 (hydrochloride) is deuterium labeled Paroxetine (hydrochloride). Paroxetine hydrochloride is a potent selective serotonin-reuptake inhibitor, commonly prescribed as an and has GRK2 inhibitory ability with IC50 of 14μM.

Cat. No.: HY-B0492S1

Purity: >98%

Clinical Data: No Development Reported

Parthenolide

((-)-Parthenolide)

Parthenolide is a sesquiterpene lactone found in the medicinal herb Feverfew. Parthenolide exhibits anti-inflammatory activity by inhibiting NF-kB activation; also inhibits HDAC1 protein without affecting other class I/II HDACs.



Cat. No.: HY-N0141

Purity: 99.13% Clinical Data: Phase 2

Size: 10 mM × 1 mL, 50 mg, 100 mg, 200 mg

Size 1 mg, 5 mg

Patulin

(Terinin) Cat. No.: HY-N6779

Patulin (Terinin) is a mycotoxin produced by fungi including the Aspergillus, Penicillium, and Byssochlamys species, is suspected to be clastogenic, mutagenic, teratogenic and cytotoxic.



99.47% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 25 mg

Pazopanib

(GW786034) Cat. No.: HY-10208

Pazopanib (GW786034) is a novel multi-target inhibitor of VEGFR1, VEGFR2, VEGFR3, PDGFRB, c-Kit, FGFR1, and c-Fms with IC_{so}s of 10, 30, 47, 84, 74, 140 and 146 nM, respectively.



99.77% Purity: Clinical Data: Launched

10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg, 500 mg

Pazopanib Hydrochloride

(GW786034 (Hydrochloride))

Pazopanib Hydrochloride (GW786034 Hydrochloride) is a novel multi-target inhibitor of VEGFR1, VEGFR2, VEGFR3, PDGFRβ, c-Kit, FGFR1, and **c-Fms** with an IC_{50} of 10, 30, 47, 84, 74, 140 and 146 nM, respectively.



Cat. No.: HY-12009

99.84% Purity: Clinical Data: Launched

10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg, 500 mg

Pazopanib-d6

(GW786034-d6) Cat. No.: HY-10208S

Pazopanib-d6 (GW786034-d6) is the deuterium labeled Pazopanib. Pazopanib (GW786034) is a novel multi-target inhibitor of VEGFR1, VEGFR2, VEGFR3, PDGFRβ, c-Kit, FGFR1, and c-Fms with IC₅₀s of 10, 30, 47, 84, 74, 140 and 146 nM, respectively.



>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

PD 169316

Cat. No.: HY-10578

PD 169316 is a potent, cell-permeable and selective p38 MAP kinase inhibitor, with IC_{so} of 89 nM. PD169316 selectively inhibits the kinase activity of the phosphorylated p38 without hindering upstream kinases to phosphorylate p38.



98.0% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 500 mg

PD-166866

Cat. No.: HY-101296

PD166866 is a selective FGFR1 tyrosine kinase inhibitor with an IC₅₀ of 52.4 nM.



99.89% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

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PD146176

(NSC168807) Cat. No.: HY-103157

PD146176 (NSC168807), a 15-Lipoxygenase (15-LO) inhibitor, inhibits rabbit reticulocyte 15-LO (K_i =197 nM, IC_{50} =0.54 μ M). PD146176 reverses cognitive impairment, brain amyloidosis, and tau pathology by stimulating autophagy in aged triple transgenic mice.



Cat. No.: HY-12028

Purity: 98 04%

PD98059

Clinical Data: No Development Reported 10 mM × 1 mL, 5 mg Size:

PD168393 is a potent, selective and

98 60%

cell-permeable inhibitor of EGFR tyrosine kinase

and ErbB2. PD168393 irreversiblely inactivates

against insulin receptor, PDGFR, FGFR and PKC.

10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 25 mg, 50 mg

EGF receptor (IC₅₀=0.7 nM) and is inactive

Clinical Data: No Development Reported

Peiminine(Verticinone; Raddeanine) is a natural compound with anti-inflammatory activity. IC50 value: Target: Peiminine and DXS significantly reduced alveolar inflammation and pulmonary interstitial inflammation in rats with bleomycin-induced lung injury.

10 mM × 1 mL, 5 mg, 10 mg

Peiminine

Purity:

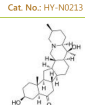
Size:

PD168393

(Verticinone; Raddeanine)

Purity:

Clinical Data: No Development Reported



Cat. No.: HY-13896

ERK1/2 signaling inhibitor. Purity:

Clinical Data: No Development Reported

PD98059 is a potent and selective MEK inhibitor

with an IC_{50} of 5 μ M. PD98059 binds to the

inactive form of MEK, thereby preventing the

activation of MEK1 (IC $_{50}$ of 2-7 μ M) and MEK2 (IC_{so} of 50 μ M) by upstream kinases. PD98059 is a

10 mM × 1 mL, 10 mg, 50 mg, 100 mg Size:

Pemetrexed

(LY231514) Cat. No.: HY-10820

Pemetrexed (LY231514) is an antifolate, the K. values of the pentaglutamate of Pemetrexed (LY231514) are 1.3, 7.2, and 65 nM for inhibits thymidylate synthase (TS), dihydrofolate reductase (DHFR), and glycinamide ribonucleotide formyltransferase (GARFT), respectively.



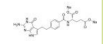
99.95% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 50 mg, 100 mg

Pemetrexed disodium

(LY231514 disodium)

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.



Cat. No.: HY-10820A

99.23% **Purity:** Clinical Data: Launched

10 mM × 1 mL, 50 mg, 100 mg, 200 mg

Pemetrexed disodium hemipenta hydrate

(LY231514 disodium hemipenta hydrate)

Pemetrexed disodium hemipenta hydrate is a novel antifolate, the K, 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.



Cat. No.: HY-13781

Purity: 99.89% Clinical Data: Launched

10 mM × 1 mL, 100 mg, 500 mg Size:

Pemetrexed-d5 disodium

(LY231514-d5 disodium)

Pemetrexed-d5 (LY231514-d5) disodium is the deuterium labeled Pemetrexed disodium.



Cat. No.: HY-10820AS

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Penfluridol

(R-16341) Cat. No.: HY-B1077

Penfluridol is a highly potent, first generation diphenylbutylpiperidine antipsychotic.

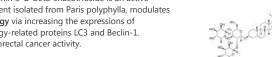


Purity: 99.93% Clinical Data: Launched

Size 10 mM × 1 mL, 50 mg, 100 mg

Pennogenin 3-O-beta-chacotrioside

Pennogenin 3-O-beta-chacotrioside is an active component isolated from Paris polyphylla, modulates autophagy via increasing the expressions of autophagy-related proteins LC3 and Beclin-1. Anti-colorectal cancer activity.



Purity: 99.93%

Clinical Data: No Development Reported 10 mM × 1 mL, 5 mg, 10 mg, 20 mg Cat. No.: HY-N4180

Pentoxifylline

(BL-191; PTX; Oxpentifylline)

Pentoxifylline (BL-191), a haemorheological agent, is an orally active non-selective **phosphodiesterase** (**PDE**) inhibitor, with immune modulation, anti-inflammatory, hemorheological, anti-fibrinolytic and anti-proliferation effects.

Cat. No.: HY-B0715

Purity: 99.35% Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 1 g

Pentoxifylline-4',4',6',6',6'-d5

Pentoxifylline-4',4',6',6',6'-d5 is the deuterium labeled Pentoxifylline.

N N O D D D

Cat. No.: HY-B0715S2

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Pentoxifylline-d6

Cat. No.: HY-B0715S

Pentoxifylline-d6 (BL-191-d6) is the deuterium labeled Pentoxifylline.

Purity: > 98%

Clinical Data: No Development Reported

Size: 1 mg, 10 mg

Pepstatin

(Pepstatin A)

Pepstatin (Pepstatin A) is a specific aspartic protease inhibitor produced by actinomycetes, with $\rm IC_{50}S$ of 4.5 nM, 6.2 nM, 150 nM, 290 nM, 520 nM and 260 nM for hemoglobin-pepsin, hemoglobin-proctase, casein-pepsin, casein-proctase, casein-acid protease...

Purity: 98.28%

Clinical Data: No Development Reported

Size: 10 mg, 50 mg



Cat. No.: HY-P0018

Peretinoin

(NIK333) Cat. No.: HY-100008

Peretinoin is an oral acyclic retinoid with a vitamin A-like structure that targets retinoid nuclear receptors such as retinoid X receptor (RXR) and retinoic acid receptor (RAR).



Purity: 99.79% Clinical Data: Launched

Size: $10 \text{ mM} \times 1 \text{ mL}, 5 \text{ mg}, 10 \text{ mg}, 50 \text{ mg}, 100 \text{ mg}$

Perifosine

(KRX-0401; NSC 639966; D21266) Cat. No.: HY-50909

Perifosine is an oral Akt inhibitor which inhibits proliferation of different tumor cell lines with IC $_{so}\text{S}$ of 0.6-8.9 $\mu\text{M}.$

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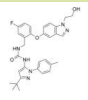
Purity: ≥98.0% Clinical Data: Phase 3

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

#### Pexmetinib

(ARRY-614) Cat. No.: HY-16782

Pexmetinib is a potent Tie-2 and p38 MAPK dual inhibitor, with  $IC_{s0}$ s of 1 nM, 35 nM and 26 nM for Tie-2, p38 $\alpha$  and p38 $\beta$ , respectively, and can be used in the research of acute myeloid leukemia.



Purity: 99.93%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

#### PF 750

PF 750 is a selective and covalent **fatty acid** 

amide hydrolase (FAAH) inhibitor, with  ${\rm IC}_{50}$ s varied from 16.2-595 nM in different pre-incubation times. Covalently modifies the enzyme's active site serine nucleophile.



**Purity:** ≥98.0%

Clinical Data: No Development Reported

Size: 1 mg

#### PF-04457845

Cat. No.: HY-14376

PF-04457845 is a highly efficacious and selective FAAH inhibitor with  $\rm IC_{so}$  values is 7.2±0.63 nM and 7.4±0.62 nM for hFAAH and rFAAH, respectively.

Purity: 99.37% Clinical Data: Phase 2

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

#### PF-04691502

PF-04691502 is a potent and selective inhibitor of PI3K and mTOR. PF-04691502 binds to human PI3K $\alpha$ ,  $\beta$ ,  $\delta$ ,  $\gamma$  and mTOR with Ks of 1.8, 2.1, 1.6, 1.9

and 16 nM, respectively.

Purity: 99.64% Clinical Data: Phase 2

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Cat. No.: HY-15177

#### PF-3845

Cat. No.: HY-14380

PF-3845 is a potent, selective, irreversible and orally active inhibitor of fatty acid amide hydrolase (FAAH), with a K<sub>i</sub> of 0.23 µM. PF-3845 is a covalent inhibitor that carbamylates FAAH's serine nucleophile.

99 90% Purity:

Clinical Data: No Development Reported

(Sphingosine Kinase 1 Inhibitor II)

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

#### PF-543

PF-543 (Sphingosine Kinase 1 Inhibitor II) is a potent, selective, reversible and sphingosine-competitive SPHK1 inhibitor with an  $\overline{IC}_{50}$  of 2 nM and a  $K_i$  of 3.6 nM. PF-543 is >100-fold selectivity for SPHK1 over SPHK2.



Cat. No.: HY-15425

Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg

#### PF-543 hydrochloride

(Sphingosine Kinase 1 Inhibitor II hydrochloride) Cat. No.: HY-15425B

PF-543 hydrochloride (Sphingosine Kinase 1 Inhibitor II hydrochloride) is a potent, selective, reversible and sphingosine-competitive SPHK1 inhibitor with an IC<sub>50</sub> of 2 nM and a K<sub>1</sub> of 3.6 nM. PF-543 hydrochloride is >100-fold selectivity for SPHK1 over SPHK2.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

### PFK-015

Cat. No.: HY-12204

PFK-015 is an effective inhibitor of PFKFB3 with IC50 of 110 nM (recombinant PFKFB3) and inhibits PFKFB3 activity in cancer cells with IC50 of 20 nM.



Purity: 98.29%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

#### PH-797804

Cat. No.: HY-10403

PH-797804 is a ATP-competitive, selective  $p38\alpha/p38\beta$  inhibitor (IC<sub>50</sub>=26 nM and K<sub>i</sub>=5.8 nM for p38α; K<sub>.</sub>=40 nM for p38β) and does not inhibit JNK2.



Purity: 98.94%

No Development Reported Clinical Data:

Size 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg, 200 mg

#### PF-4708671

PF-4708671 is a potent cell-permeable S6K1 inhibitor with a K, of 20 nM and IC, of 160 nM.



Cat. No.: HY-15773

99 94% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 10 mg, 25 mg, 50 mg, 100 mg

#### PF-543 Citrate

(Sphingosine Kinase 1 Inhibitor II Citrate)

PF-543 Citrate (Sphingosine Kinase 1 Inhibitor II Citrate) is a potent, selective, reversible and sphingosine-competitive SPHK1 inhibitor with an IC<sub>50</sub> of 2 nM and a K<sub>i</sub> of 3.6 nM. PF-543 Citrate is >100-fold selectivity for SPHK1 over SPHK2.



Cat. No.: HY-15425A

**Purity:** 98.35%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg Size:

#### PFI-1

PFI-1 is a selective BET (bromodomain-containing protein) inhibitor for BRD4 with  $IC_{50}$  of 0.22  $\mu M$ 

in a cell-free assay.

Cat. No.: HY-16586

99.88% Purity:

Clinical Data: No Development Reported

Size 10 mM × 1 mL, 10 mg, 50 mg, 100 mg

#### PFK-158

Cat. No.: HY-12203

PFK-158 is a potent and selective PFKFB3 inhibitor with an  $IC_{50}$  value 137 nM. PFK-158 reduces glucose uptake, ATP production, lactate release, and induces apoptosis and autophagy in cancer cells. PFK-158 has broad anti-tumor activity.

>98% Purity: Clinical Data: Phase 1 Size: 5 mg, 10 mg



### PHA-665752

Cat. No.: HY-11107

PHA-665752 is a selective, ATP-competitive, and active-site inhibitor of the catalytic activity of **c-Met** kinase (**K**<sub>i</sub>=4 nM; **IC**<sub>so</sub>=9 nM). PHA-665752 exhibits > 50-fold selectivity for c-Met compared with a panel of diverse tyrosine and serine-threonine kinases.



Purity: 99.85%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

#### Phenformin hydrochloride

(Phenethylbiguanide hydrochloride)

Phenformin hydrochloride is an anti-diabetic drug from the biguanide class, can activate AMPK activity.

Cat. No.: HY-16397A

98 12% Purity: Clinical Data: Launched

Size:  $10 \text{ mM} \times 1 \text{ mL}, 500 \text{ mg}, 1 \text{ g}, 5 \text{ g}$ 

#### Phenylbutyrate-d11 sodium (4-PBA-d11 sodium; 4-Phenylbutyric

acid-d11 sodium; Benzenebutyric acid-d11 sodium)

Phenylbutyrate-d11 (sodium) is deuterium labeled Sodium 4-phenylbutyrate, Sodium 4-phenylbutyrate (4-PBA sodium) is an inhibitor of HDAC and endoplasmic reticulum (ER) stress, used in cancer and infection research.

Cat. No.: HY-15654S

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Physalin A

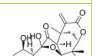
Cat. No.: HY-N9942

Physalin A is a withanolide isolated from Physalis alkekengi var. franchetii. Physalin A induces apoptosis associated with up-regulation of caspase-3 and caspase-8 expression. Physalin A induces autophagy, found to antagonize apoptosis in HT1080 cells.

Purity:

Clinical Data: No Development Reported

1 mg, 5 mg



Size:

## Cat. No.: HY-10115

PI-103 is a potent PI3K and mTOR inhibitor with IC<sub>50</sub>s of 8 nM, 88 nM, 48 nM, 150 nM, 20 nM, and 83 nM for p110α, p110β, p110δ, p110γ, mTORC1, and mTORC2. PI-103 also inhibits DNA-PK with an IC50 of 2 nM. PI-103 induces autophagy.

Purity: 98.93%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

### Physalin B

Physalin B, one of the major active steroidal constituents of Cape gooseberry, induces cell cycle arrest and triggers apoptosis in breast cancer cells through modulating p53-dependent apoptotic pathway.

**Purity:** 96.90%

Clinical Data: No Development Reported



Cat. No.: HY-N7695

## PI-103



#### PI-103 Hydrochloride

Cat. No.: HY-10115A

PI-103 Hydrochloride is a dual PI3K and mTOR inhibitor with  $IC_{50}$ s of 8 nM, 88 nM, 48 nM, 150 nM, 20 nM, and 83 nM for p110α, p110β, p110δ, p110y, mTORC1, and mTORC2. PI-103 Hydrochloride also inhibits DNA-PK with an IC50 of 2 nM. PI-103 Hydrochloride induces autophagy.

98.06% **Purity:** 

Clinical Data: No Development Reported

Size 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

#### PI-103-d8

#### Cat. No.: HY-10115S

PI-103-d8 is the deuterium labeled PI-103. PI-103 is a potent PI3K and mTOR inhibitor with IC<sub>so</sub>s of 8 nM, 88 nM, 48 nM, 150 nM, 20 nM, and 83 nM for p110 $\alpha$ , p110 $\beta$ , p110 $\delta$ , p110 $\gamma$ , mTORC1, and mTORC2. PI-103 also inhibits DNA-PK with an IC50 of 2 nM. PI-103 induces autophagy.

Purity:

Clinical Data: No Development Reported

Size 1 mg, 5 mg



#### Piceatannol

#### (Astringenin; trans-Piceatannol)

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-13518

Purity: 98.09%

Clinical Data: No Development Reported

10 mM × 1 mL, 10 mg, 25 mg, 50 mg, 100 mg Size:

## **Pictilisib**

#### (GDC-0941) Cat. No.: HY-50094

Pictilisib (GDC-0941) is a potent inhibitor of PI3Kα/δ with an IC<sub>50</sub> of 3 nM, with modest selectivity against p110β (11-fold) and p110γ (25-fold).



Purity: 99.80% Clinical Data: Phase 2

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg

#### Pictilisib dimethanesulfonate

(GDC-0941 dimethanesulfonate; GDC-0941 2 MeSO3H salt) Cat. No.: HY-20180

Pictilisib dimethanesulfonate (GDC-0941 dimethanesulfonate) is a potent inhibitor of PI3Kα/δ with IC<sub>50</sub> of 3 nM, with modest selectivity against p110 $\beta$  (11-fold) and p110 $\gamma$  (25-fold).



Purity: 99.31% Clinical Data: Phase 2

10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg

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#### Pifithrin-µ

(PFTµ; 2-Phenylethynesulfonamide)

Cat. No.: HY-10940

((+)-Pinocoembrin; Dihydrochrysin; Galangin flavanone)

Cat. No.: HY-N0575

Pifithrin-µ is an inhibitor of p53 and HSP70, with antitumor and neuroprotective activity.

98 31% Purity:

Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg

Pinocembrin ((+)-Pinocoembrin) is a flavonoid found in propolis, acts as a competitive inhibitor of histidine decarboxylase, and is an effective anti-allergic agent, with antioxidant, antimicrobial and anti-inflammatory properties.

Purity: 99.65%

Pinocembrin

10 mM × 1 mL, 5 mg, 10 mg, 25 mg Size:

Clinical Data: No Development Reported

#### Pinosylvin

Cat. No.: HY-N2387

Pinosylvin is a pre-infectious stilbenoid toxin isolated from the heartwood of Pinus spp, has anti-bacterial activities. Pinosylvin is a resveratrol analogue, can induce cell apoptosis and autophapy in leukemia cells.

Purity: 99.66%

Clinical Data: No Development Reported

5 mg, 10 mg Size:

#### **Piperine**

(Bioperine; 1-Piperoylpiperidine)

Piperine, a natural alkaloid isolated from Piper nigrum L, inhibits P-glycoprotein and CYP3A4 activities with an  $IC_{50}$  value of  $61.94\pm0.054~\mu g/mL$ in HeLa cell.



Cat. No.: HY-N0144

**Purity:** 98 88% Clinical Data: Phase 2

10 mM × 1 mL, 200 mg, 1 g, 5 g

Piperlongumine

Purity:

(Piplartine) Cat. No.: HY-N2329

Piperlongumine is a alkaloid, possesses ant-inflammatory, antibacterial, antiangiogenic, antioxidant, antitumor, and antidiabetic activities. Piperlongumine induces ROS, and induces apoptosis in cancer cell lines.



Pirarubicin (THP)

Cat. No.: HY-13725

Pirarubicin is an anthracycline antibiotics, acts as a topoisomerase II inhibitor, and is a widely used for treatment of various cancers, in particular, solid tumors.



Purity: 99.61% Clinical Data: Launched

Size 10 mg, 50 mg, 100 mg

Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg

99.19%

#### Pirarubicin Hydrochloride

(THP Hydrochloride) Cat. No.: HY-13725A

Pirarubicin Hydrochloride is an anthracycline antibiotics, acts as a topoisomerase II inhibitor, and is a widely used for treatment of various cancers, in particular, solid tumors.

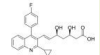
98.51% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg

#### Pitavastatin

(NK-104) Cat. No.: HY-B0144A

Pitavastatin (NK-104) is a potent hydroxymethylglutaryl-CoA (HMG-CoA) reductase inhibitor. Pitavastatin inhibits cholesterol synthesis from acetic acid with an IC<sub>so</sub> of 5.8 nM in HepG2 cells.



Purity: >98% Clinical Data: Launched Size: 1 mg, 5 mg

#### Pitavastatin Calcium

(NK-104 hemicalcium; Pitavastatin hemicalcium) Cat. No.: HY-B0144

Pitavastatin Calcium (NK-104 hemicalcium) is a potent hydroxymethylglutaryl-CoA (HMG-CoA) reductase inhibitor. Pitavastatin Calcium (NK-104 hemicalcium) inhibits cholesterol synthesis from acetic acid with an IC<sub>50</sub> of 5.8 nM in HepG2 cells.



Purity: 99.45% Clinical Data: Launched

Size 10 mM × 1 mL, 10 mg, 50 mg, 100 mg

## Pitavastatin D4

(NK-104 D4) Cat. No.: HY-B0144AS

Pitavastatin D4 (NK-104 D4) is deuterium labeled Pitavastatin. Pitavastatin is a potent HMG-CoA reductase inhibitor.



Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg

#### Pitavastatin-d4 hemicalcium

(NK-104-d4 hemicalcium; Pitavastatin-d4 hemicalcium) Cat. No.: HY-B0144S

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

#### Poloppin Polydatin

Poloppin is a potent, cell penetrant inhibitor of the mitotic Polo-like kinase (PLK) ( $IC_{so}$ =26.9  $\mu$ M) and prevents the protein-protein interaction via the Polo-box domain (PBD) ( $K_d$ = 29.5  $\mu$ M).

Purity: >98%

Clinical Data: No Development Reported

5 mg, 10 mg Size:

## Cat. No.: HY-124761 (Piceid)

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.

Pogostone is isolated from patchouli with

99 80%

Clinical Data: No Development Reported

anti-bacterial and anti-cancer

98 55% **Purity:** Clinical Data: Phase 2

10 mM × 1 mL, 50 mg, 100 mg, 200 mg, 500 mg

5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Polyphyllin G

#### Cat. No.: HY-N0817

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).



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

## Polyphyllin I

Pogostone

activities.

**Purity:** 

Size:

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.

99.61% **Purity:** 

Clinical Data: No Development Reported Size 5 mg, 10 mg, 20 mg



#### (AP24534) Cat. No.: HY-12047

Ponatinib (AP24534) is an orally active multi-targeted kinase inhibitor with IC<sub>50</sub>s of 0.37 nM, 1.1 nM, 1.5 nM, 2.2 nM, and 5.4 nM for Abl, PDGFRα, VEGFR2, FGFR1, and Src, respectively.



99.43% Purity: Clinical Data: Launched

Size 10 mM × 1 mL, 10 mg, 50 mg, 100 mg

#### Ponatinib hydrochloride (AP24534 hydrochloride)

Ponatinib (AP24534) hydrochloride is a hydrochloride of ponatinib. Ponatinib is an orally active multi-targeted kinase inhibitor with IC<sub>so</sub>s of 0.37 nM, 1.1 nM, 1.5 nM, 2.2 nM, and 5.4 nM for Abl, PDGFRa, VEGFR2, FGFR1, and Src, respectively.

Purity: >98% Clinical Data: Launched 1 mg, 5 mg Size:



Cat. No.: HY-108766

Cat. No.: HY-N1416

Cat. No.: HY-N0120A

Cat. No.: HY-N0047

#### Ponatinib-d8

#### (AP24534-d8) Cat. No.: HY-12047S

Ponatinib D8 (AP24534 D8) is a deuterium labeled Ponatinib. Ponatinib (AP24534) is an orally active multi-targeted kinase inhibitor with IC<sub>sos</sub> of 0.37 nM, 1.1 nM, 1.5 nM, 2.2 nM, and 5.4 nM for Abl, PDGFRα, VEGFR2, FGFR1, and Src, respectively.



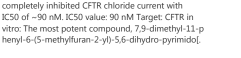
Purity: 98.44%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

### PPQ-102 (CFTR Inhibitor)

PPQ-102 is a potent CFTR inhibitor which can completely inhibited CFTR chloride current with IC50 of ~90 nM. IC50 value: 90 nM Target: CFTR in vitro: The most potent compound, 7,9-dimethyl-11-p



Purity: 99.82%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Cat. No.: HY-14179

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

#### PR-619

PR-619 is a broad-range and reversible DUB

inhibitor with EC<sub>so</sub>s of 3.93, 4.9, 6.86, 7.2, and 8.61 µM for USP4, USP8, USP7, USP2, and USP5, respectively. PR-619 induces ER Stress and ER-Stress related apoptosis.

Purity: 98 89%

Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 50 mg

#### Cat. No.: HY-13814

## Pravastatin sodium

(CS-514 sodium) Cat. No.: HY-B0165A

Pravastatin sodium (CS-514 sodium) is an HMG-CoA reductase inhibitor against sterol synthesis with  $IC_{50}$  of 5.6  $\mu$ M.

Purity: 99 49% Clinical Data: Launched

10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg Size:

## Pravastatin

(CS-514)Cat. No.: HY-B0165

Pravastatin (CS-514) is a competitive HMG-CoA reductase inhibitor against sterol synthesis with  $IC_{50}$  of 5.6  $\mu$ M.



**Purity:** >98% Clinical Data: Launched Size: 5 mg, 10 mg

#### Pravastatin-13C,d3 sodium

(CS-514-13C,d3 sodium)

Pravastatin-13C,d3 (sodium) is the 13C- and deuterium labeled, Pravastatin sodium (CS-514 sodium) is an HMG-CoA reductase inhibitor against sterol synthesis with IC50 of 5.6  $\mu$ M.



Cat. No.: HY-B0165AS

**Purity:** >98%

Clinical Data: No Development Reported

1 mg, 5 mg

#### Pravastatin-d3 sodium salt

Cat. No.: HY-B0165CS

Pravastatin-d3 (CS-514-d3) sodium salt is the deuterium labeled Pravastatin sodium salt. Pravastatin (CS-514) sodium salt is a competitive HMG-CoA reductase inhibitor against sterol synthesis with  $IC_{so}$  of 5.6  $\mu$ M.



>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

## Prazosin hydrochloride

Cat. No.: HY-B0193A

Prazosin hydrochloride is a well-tolerated, CNS-active  $\alpha 1$ -adrenergic receptor antagonist for the research of high blood pressure and alcohol use disorders.



**Purity:** 99 93% Clinical Data: Launched

Size  $10~\text{mM}\times1~\text{mL},\,100~\text{mg},\,200~\text{mg},\,500~\text{mg}$ 

#### Pregnenolone

(3β-Hydroxy-5-pregnen-20-one)

Pregnenolone (3β-Hydroxy-5-pregnen-20-one) is a powerful neurosteroid, the main precursor of various steroid hormones including steroid ketones



Cat. No.: HY-B0151

98.05% Purity: Clinical Data: Launched

10 mM × 1 mL, 500 mg, 1 g, 5 g Size:

#### Pregnenolone monosulfate

(3β-Hydroxy-5-pregnen-20-one monosulfate)

Pregnenolone monosulfate (3β-Hydroxy-5-pregnen-20-one monosulfate) is a powerful neurosteroid, the main precursor of various steroid hormones including steroid ketones



Cat. No.: HY-B1739

≥98.0% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg

#### Pregnenolone monosulfate sodium

(3β-Hydroxy-5-pregnen-20-one monosulfate sodium) Cat. No.: HY-110189

Pregnenolone monosulfate sodium (3β-Hydroxy-5-pregnen-20-one monosulfate sodium) is a powerful neurosteroid, the main precursor of various steroid hormones including steroid ketones.



Purity: ≥95.0% Clinical Data: Launched

5 mg, 10 mg, 50 mg, 100 mg

#### Pregnenolone monosulfate-d4 sodium

(3β-Hydroxy-5-pregnen-20-one monosulfate-d4 sodium) Cat. No.: HY-110189S1

Pregnenolone monosulfate-d4 (sodium) is the deuterium labeled Pregnenolone monosulfate.



Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg

#### Pregnenolone-d4-1

 $(3\beta$ -Hydroxy-5-pregnen-20-one-d4-1)

Pregnenolone-d4-1 (3B-Hvdroxy-5-preanen-20-one-d4-1) is the

Cat. No.: HY-B0151S2

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

deuterium labeled Pregnenolone.

#### PRIMA-1

(NSC-281668)

PRIMA-1 (NSC-281668) is a mutant p53 reactivator, restores the sensitivity of TP53 mutant-type thyroid cancer cells to the histone methylation inhibitor 3-Deazaneplanocin A.

OH

Cat. No.: HY-19980A

>98.0% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg, 200 mg

### Procainamide hydrochloride

Cat. No.: HY-A0084

Procainamide hydrochloride is an anti-arrhythmic agent and is used to treat cardiac arrhythmia; induces rapid block of the batrachotoxin(BTX)-activated sodium channels of the heart muscle and acts as antagonist to long gating closures.

≥98.0% Purity: Clinical Data: Launched

10 mM × 1 mL, 100 mg

#### Proflavine hemisulfate

(Proflavin hemisulfate; 3,6-Diaminoacridine hemisulfate)

Proflavine hemisulfate, an acridine dye, is a known DNA intercalating agent. Anti-microbial agent. Proflavine hemisulfate behaves as a pore blocker for K, 3.2. Proflavine hemisulfate is a potential lead compound for K<sub>ir</sub>3.2-associated neurological diseases.

98.17% **Purity:** Clinical Data: Phase 2

10 mM × 1 mL, 100 mg



Cat. No.: HY-B0883

0.5H<sub>2</sub>SO<sub>4</sub>

Cat. No.: HY-15888

#### Pseudolaric Acid B

Cat. No.: HY-N6939

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: 99.47%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

#### PTC-209

PTC-209 is a specific BMI-1 inhibitor with an IC<sub>so</sub> of 0.5 μM in HEK293T cell line. PTC-209 irreversibly impairs colorectal cancer-initiating cells (CICs). PTC-209 shows potent anti-myeloma

activity and impairs the tumor microenvironment.

99.85% Purity:

Clinical Data: No Development Reported

Size 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

### PTC-209 hydrobromide

Cat. No.: HY-15888A

PTC-209 hydrobromide is a specific **BMI-1** inhibitor with an  $IC_{50}$  of 0.5  $\mu M$  in HEK293T cell line. PTC-209 hydrobromide irreversibly impairs colorectal cancer-initiating cells (CICs). PTC-209 hydrobromide shows potent anti-myeloma activity and impairs the tumor microenvironment.

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg



## Pterostilbene

Pterostilbene is a stilbenoid isolated from blueberries and Pterocarpus marsupium. Shows anti-oxidant, anti-inflammatory, anti-carcinogenic, anti-diabetic and anti-obesity properties.

Purity: 99.79% Clinical Data: Phase 3

10 mM × 1 mL, 25 mg Size:



Cat. No.: HY-N0828

#### Purvalanol A

(NG-60) Cat. No.: HY-18299A

Purvalanol A is a potent CDK inhibitor, which inhibits cdc2-cyclin B, cdk2-cyclin A, cdk2-cyclin E, cdk4-cyclin D1, and cdk5-p35 with ICsos of 4, 70, 35, 850, 75 nM, resepctively.



99.11%

Clinical Data: No Development Reported

10 mM × 1 mL, 10 mg, 25 mg, 50 mg, 100 mg

## Purmorphamine

Cat. No.: HY-15108

Purmorphamine is a smoothened/Smo receptor agonist with an  $EC_{50}$  of 1  $\mu$ M.

Purity: 99.89%

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

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#### PX-478

Cat. No.: HY-10231

PX-478 is an orally active HIF- $1\alpha$  inhibitor with potent antitumor activities. PX-478 can cross the blood-brain barrier.

HCI

Cat. No.: HY-B0271

Purity: > 98.0% Clinical Data: Phase 1

Size: 5 mg, 10 mg, 50 mg, 100 mg

#### PX20606 trans racemate

(PX-102 trans racemate)

PX20606 trans racemate (PX-102 trans racemate) is a FXR agonist with EC<sub>so</sub>s of 32 and 34 nM for FXR in FRET and M1H assay, respectively.



Cat. No.: HY-100443A

99.01% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 1 mg, 2 mg

#### Pyrazinamide

(Pyrazinecarboxamide; Pyrazinoic acid amide)

Pyrazinamide (Pyrazinecarboxamide; Pyrazinoic acid amide) is a potent and orally active antitubercular antibiotic. Pyrazinamide is a prodrug that is converted to the active form pyrazinoic acid (POA) by PZase/nicotinamidase encoded by the pncA gene in M. tuberculosis.

Purity: Clinical Data: Launched

10 mM × 1 mL, 500 mg, 10 g, 50 g

#### Pyrazinamide-d3

(Pyrazinecarboxamide-d3; Pyrazinoic acid amide-d3)

Pyrazinamide-d3 is deuterium labeled Pyrazinamide. Pyrazinamide (Pyrazinecarboxamide; Pyrazinoic acid amide) is a potent and orally active antitubercular antibiotic.

Cat. No.: HY-B0271S

**Purity:** >98%

Clinical Data: No Development Reported

1 mg, 5 mg

#### Pyriproxyfen

(S-31183) Cat. No.: HY-B2031

Pyriproxyfen is a juvenile hormone analog, preventing larvae from developing into adulthood and thus rendering them unable to reproduce. Pyriproxyfen is a pyridine-based pesticide which is found to be effective against a variety of arthropoda.

Purity: 99.70%

Clinical Data: No Development Reported Size: 10 mM × 1 mL, 500 ma, 5 a

#### Quercetin

Quercetin, a natural flavonoid, is a stimulator of recombinant SIRT1 and also a PI3K inhibitor with  $IC_{so}$  of 2.4  $\mu$ M, 3.0  $\mu$ M and 5.4  $\mu$ M for PI3K  $\gamma$ , PI3K

δ and PI3K β, respectively.

Cat. No.: HY-18085

98 02% Purity: Clinical Data: Phase 4

Size 10 mM  $\times$  1 mL, 500 mg, 1 g, 5 g

### Quercetin-d3

Cat. No.: HY-18085S1

Quercetin-d3 is the deuterium labeled Quercetin. Quercetin, a natural flavonoid, is a stimulator of recombinant SIRT1 and also a PI3K inhibitor with  $IC_{so}$  of 2.4  $\mu$ M, 3.0  $\mu$ M and 5.4  $\mu$ M for PI3K  $\gamma$ , PI3K δ and PI3K β, respectively.

>98% **Purity:** 

Quercitrin

Clinical Data: No Development Reported

2.5 mg, 25 mg Size

(Quercetin 3-rhamnoside) Cat. No.: HY-N0418

Quercitrin is a natural compound found in Tartary buckwheat with a potential anti-inflammation effect that is used to treat heart and vascular conditions.

Purity: 99.80%

No Development Reported Clinical Data:

Size 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

#### Quercetin-d5

Quercetin-d5 is a deuterium labeled Quercetin. Quercetin, a natural flavonoid, is a stimulator of recombinant SIRT1 and also a PI3K inhibitor with  $IC_{so}$  of 2.4  $\mu$ M, 3.0  $\mu$ M and 5.4  $\mu$ M for PI3K  $\gamma$ , PI3K δ and PI3K β, respectively.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

Cat. No.: HY-18085S

#### Quinacrine dihydrochloride

(Mepacrine dihydrochloride; SN-390 dihydrochloride) Cat. No.: HY-13735A

Quinacrine (Mepacrine) dihydrochloride is an orally bioavailable antimalarial agent, which possess anticancer effect both in vitro and vivo. Quinacrine dihydrochloride suppresses NF-κB and activate p53 signaling, which results in the induction of the apoptosis.

Purity: 99.01% Clinical Data: Phase 2

10 mM × 1 mL, 100 mg, 500 mg



#### Quinacrine hydrochloride hydrate (Mepacrine hydrochloride

hydrate; SN-390 hydrochloride hydrate) Cat. No.: HY-13735B

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.

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

### Quizartinib

Purity:

Size:

Quisinostat

(JNJ-26481585)

antitumoral activity.

Clinical Data: Phase 2

Quisinostat (JNJ-26481585) is a potent,

98.02%

second-generation and orally active pan-HDAC

inhibitor (HDACi), with  $IC_{50}$  values ranging from

and HDAC11. Quisinostat has a broad spectrum

0.11 nM to 0.64 nM for HDAC1, HDAC2, HDAC4, HDAC10

(AC220) Cat. No.: HY-13001

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Quizartinib (AC220) is an orally active, highly selective and potent second-generation type II FLT3 tyrosine kinase inhibitor, with a K<sub>d</sub> of 1.6 nM. Quizartinib inhibits wild-type FLT3 and FLT3-ITD autophosphorylation in MV4-11 cells with

IC<sub>so</sub>s of 4.2 and 1.1 nM, respectively.

**Purity:** 99.01% Clinical Data: Launched

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

Quisinostat dihydrochloride

(JNJ-26481585 dihydrochloride) Cat. No.: HY-15433A

Quisinostat dihydrochloride (JNJ-26481585 dihydrochloride) is an orally available, potent pan-HDAC inhibitor with IC<sub>so</sub>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.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

QW24

Cat. No.: HY-143655

QW24 exerts potent anti-tumor activity by down-regulating BMI-1 and is used as an effective therapeutic agent for clinical colorectal cancer treatment.

>98% Purity:

Clinical Data: No Development Reported

Size: 1 ma, 5 ma QX77

QX77 is a chaperone-mediated autophagy (CMA) activator and upregulates LAMP2A expression in vitro. QX77 induces Rab11 upregulation, rescues Rab11 down-regulation and trafficking deficiency in cystinotic cells. QX77 can impede self-renewal and promote differentiation of ES cells.

**Purity:** 99.48%

Clinical Data: No Development Reported 5 mg, 10 mg, 50 mg, 100 mg Size:

R1487 Hydrochloride

Cat. No.: HY-14975

R1487 Hydrochloride is a highly potent and selective  $p38\alpha$  inhibitor, with  $K_a$  values of 0.2 nM and 29 nM for p38α and p38β, respectively.

98.94% Purity:

Clinical Data: No Development Reported

Size 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg Rabusertib

(LY2603618; IC-83) Cat. No.: HY-14720

Rabusertib (LY2603618) is a potent and selective inhibitor of Chk1 with an IC<sub>50</sub> of 7 nM.

Cat. No.: HY-112483

Cat. No.: HY-15433

99.73% Purity: Clinical Data: Phase 2

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

RAF265

(CHIR-265) Cat. No.: HY-10248

RAF265 is a potent RAF/VEGFR2 inhibitor.

99.98% Purity: Clinical Data: Phase 2

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg Size:

Ralimetinib

(LY2228820) Cat. No.: HY-13241A

Ralimetinib (LY2228820) is a potent and selective, ATP-competitive inhibitor of p38 MAPK  $\alpha/\beta$ , with IC<sub>so</sub>s of 5.3 and 3.2 nM, respectively. Ralimetinib (LY2228820) selectively inhibits phosphorylation of MK2 (Thr334), with no effect on phosphorylation of p38α MAPK, JNK, ERK1/2, c-Jun, ATF2, or c-Myc.

>98%

Clinical Data: No Development Reported

1 mg, 5 mg

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#### Ralimetinib dimesylate

(LY2228820 dimesylate) Cat. No.: HY-13241

Ralimetinib dimesylate (LY2228820 dimesylate) is a selective. ATP-competitive inhibitor of p38 MAPK  $\alpha/\beta$  with IC<sub>50</sub>s of 5.3 and 3.2 nM, respectively.



99 52% Purity: Clinical Data: Phase 2

Size:  $10 \text{ mM} \times 1 \text{ mL}$ , 5 mg, 10 mg, 50 mg, 100 mg

#### Raloxifene hydrochloride (Keoxifene hydrochloride; LY156758;

LY139481 hydrochloride) Cat. No.: HY-13738A

Raloxifene hydrochloride (Keoxifene hydrochloride) is a second generation selective and orally active estrogen receptor modulator.



Purity: 99 91% Clinical Data: Launched

Size: 10 mM × 1 mL, 50 mg, 100 mg

#### Ranolazine dihydrochloride

(CVT 303 dihydrochloride; RS 43285) Cat. No.: HY-17401

Ranolazine dihydrochloride (CVT 303 dihydrochloride) is an anti-angina drug that achieves its effects by inhibiting the late phase of inward sodium current ( $I_{Na}$  and  $I_{Kr}$  with  $IC_{50}$ values of 6 μM and 12 μM, respectively) without affecting heart rate or blood pressure...



Purity: Clinical Data: Launched

10 mM × 1 mL, 100 mg, 200 mg, 500 mg, 1 g, 5 g Size:

#### Ranolazine-d8 dihydrochloride

(CVT 303-d8 dihydrochloride; RS 43285-d8)

Ranolazine-d8 (CVT 303-d8) dihydrochloride is the deuterium labeled Ranolazine dihydrochloride.



Cat. No.: HY-17401S

**Purity:** >98%

Clinical Data: No Development Reported

1 mg, 5 mg

#### RapaLink-1

Cat. No.: HY-111373

RapaLink-1, the third-generation bivalent mTOR inhibitor, combines Rapamycin (HY-10219) with MLN0128 (HY-13328, a second-generation mTOR kinase inhibitor) by an inert chemical linker.



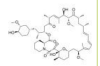
97.93% Purity:

Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg, 25 mg

#### Rapamycin

(Sirolimus; AY-22989)

Rapamycin (Sirolimus; AY 22989) is a potent and specific mTOR inhibitor with an IC<sub>50</sub> of 0.1 nM in HEK293 cells. Rapamycin binds to FKBP12 and specifically acts as an allosteric inhibitor of mTORC1. Rapamycin is an autophagy activator, an immunosuppressant.



Cat. No.: HY-10219

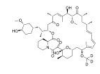
99.94% **Purity:** Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 25 mg, 50 mg, 100 mg, 200 mg

#### Rapamycin-d3

(Sirolimus-d3; AY-22989-d3) Cat. No.: HY-10219S

Rapamycin-d3 (Sirolimus-d3) is the deuterium labeled Rapamycin. Rapamycin is a potent and specific mTOR inhibitor with an IC<sub>so</sub>of 0.1 nM in HEK293 cells. Rapamycin binds to FKBP12 and specifically acts as an allosteric inhibitor of mTORC1.



Purity: 95.30%

Clinical Data: No Development Reported Size 1 mg, 5 mg, 10 mg

#### Rasagiline mesylate

((R)-AGN1135 mesylate; TVP1012 mesylate)

Rasagiline (R-AGN1135) mesylate is a highly potent selective irreversible mitochondrial monoamine oxidase (MAO) inhibitor with IC<sub>so</sub>s of 4.43nM and 412nM for rat brain MAO B and A activity, respectively.



Cat. No.: HY-14605

99.66% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 50 mg, 100 mg

## Regorafenib

(BAY 73-4506) Cat. No.: HY-10331

Regorafenib (BAY 73-4506) is a multi-targeted receptor tyrosine kinase inhibitor with ICsos of 13/4.2/46, 22, 7, 1.5 and 2.5 nM for VEGFR1/2/3, PDGFRβ, Kit, RET and Raf-1, respectively.



Purity: 99.65% Clinical Data: Launched

10 mM × 1 mL, 10 mg, 50 mg, 100 mg Size:

## Regorafenib Hydrochloride

(BAY 73-4506 hydrochloride)

Regorafenib Hydrochloride (BAY 73-4506 hydrochloride) is a multi-target inhibitor for VEGFR1/2/3, PDGFRβ, Kit, RET and Raf-1 with IC<sub>50</sub>s of 13/4.2/46, 22, 7, 1.5 and 2.5 nM, respectively.



Cat. No.: HY-13308

Purity: 99.58% Clinical Data: Launched

10 mM × 1 mL, 10 mg, 50 mg, 100 mg

#### Regorafenib monohydrate

(BAY 73-4506 monohydrate) Cat. No.: HY-10331A

Regorafenib monohydrate (BAY 73-4506 monohydrate) is a multi-target inhibitor for VEGFR1/2/3, PDGFR $\beta$ , Kit, RET and Raf-1 with IC  $_{so}$ S of 13/4.2/46, 22, 7, 1.5 and 2.5 nM, respectively.

Purity: 99.96% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg

## Regorafenib-13C,d3

(BAY 73-4506-13C,d3)

Regorafenib-13C,d3 is the 13C- and deuterium labeled. Regorafenib (BAY 73-4506) is a multi-targeted receptor tyrosine kinase inhibitor with IC50s of 13/4.2/46, 22, 7, 1.5 and 2.5 nM for VEGFR1/2/3, PDGFRβ, Kit, RET and Raf-1, respectively.



Cat. No.: HY-10331S1

**Purity:** >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Regorafenib-d3

(BAY 73-4506-d3) Cat. No.: HY-10331S

Regorafenib D3 (BAY 73-4506 D3) is a deuterium labeled Regorafenib. Regorafenib is a multi-targeted receptor tyrosine kinase inhibitor.

**Purity:** >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Resatorvid

(TAK-242; CLI-095)

Resatorvid (TAK-242) is a selective **Toll-like receptor 4** (TLR4) inhibitor. Resatorvid inhibits NO, TNF- $\alpha$  and IL-6 production with IC $_{50s}$  of 1.8 nM, 1.9 nM and 1.3 nM, respectively. Resatorvid downregulates expression of TLR4 downstream signaling molecules MyD88 and TRIF.

Purity: 99.95% Clinical Data: Phase 3

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

HŅ D=S=O O

Cat. No.: HY-11109

#### Reserpine

Cat. No.: HY-N0480

Reserpine is an inhibitor of the **vesicular** monoamine transporter 2 (VMAT2).



Purity: 99.83% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg

### Reserpine-d9

Reserpine-d9 is the deuterium labeled Reserpine. Reserpine is an inhibitor of the **vesicular monoamine transporter 2 (VMAT2)**.



Cat. No.: HY-N0480S

**Purity:** >98%

Clinical Data: No Development Reported

**Size:** 2.5 mg, 25 mg

#### Resveratrol

(trans-Resveratrol; SRT501) Cat. No.: HY-16561

Resveratrol (trans-Resveratrol; SRT501), a natural polyphenolic phytoalexin that possesses anti-oxidant, anti-inflammatory, cardioprotective, and anti-cancer properties.

Purity: 99.89%
Clinical Data: Launched

Size: 10 mM × 1 mL, 200 mg, 500 mg

#### Resveratrol analog 1

Resveratrol analog 1 is an analog of Resveratrol (HY-16561), compound 48. Resveratrol is a natural polyphenolic phytoalexin that possesses anti-oxidant, anti-inflammatory, cardioprotective, and anti-cancer properties.



Cat. No.: HY-136203

**Purity:** 98.06%

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

#### Resveratrol analog 2

Cat. No.: HY-136204

Resveratrol analog 2 is an **analog** of Resveratrol (HY-16561). Resveratrol is a natural polyphenolic phytoalexin that possesses anti-oxidant, anti-inflammatory, cardioprotective, and anti-cancer properties.



**Purity:** > 98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Resveratrol-d4

(trans-Resveratrol-d4; SRT501-d4)

Resveratrol-d4 (trans-Resveratrol-d4) is the deuterium labeled Resveratrol. Resveratrol (trans-Resveratrol; SRT501), a natural polyphenolic phytoalexin that possesses anti-oxidant, anti-inflammatory, cardioprotective, and anti-cancer properties.



Cat. No.: HY-16561S

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

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#### Retinoic acid

(Vitamin A acid; all-trans-Retinoic acid; ATRA)

Retinoic acid is a metabolite of vitamin A that plays important roles in cell growth. differentiation, and organogenesis. Retinoic acid is a natural agonist of RAR nuclear receptors, with  $IC_{so}$ s of 14 nM for RAR $\alpha/\beta/\gamma$ . Retinoic acid bind to PPAR $\beta/\delta$  with  $K_d$  of 17 nM.

Cat. No.: HY-14649

Purity: 99 74% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg, 1 g, 5 g

#### Retro-2

Retro-2 is a selective inhibitor of retrograde protein trafficking at the endosome-trans-Golgi network interface. Retro-2 is an ebolavirus (EBOV) infection inhibitor with an EC<sub>50</sub> of 12.2 μM in HeLa cells. Retro-2 induces cell autophagy.



Cat. No.: HY-N0105

Cat. No.: HY-122571

Purity: ≥98.0%

Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

## Reversine

Cat. No.: HY-14711

Reversine is a novel class of ATP-competitive Aurora kinase inhibitor with IC<sub>50</sub>s of 400, 500 and 400 nM for Aurora A, Aurora B and Aurora C, respectively.

Purity: 99 40%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

#### Rhein

(Rheic Acid; Rhubarb yellow; Monorhein)

Rhein is a lipophilic anthraguinone extensively found in medicinal herbs, and has many pharmacological effects, including epatoprotective, nephroprotective, anti-inflammatory, antioxidant, anticancer, and antimicrobial activities.

**Purity:** 

Clinical Data: No Development Reported

10 mM × 1 mL, 100 mg, 200 mg, 500 mg

### Rhoifolin

Cat. No.: HY-N0755

Rhoifolin is a flavone glycoside isolated from Citrus grandis (L.) Osbeck leaves. Rhoifolin is beneficial for diabetic complications through enhanced adiponectin secretion, tyrosine phosphorylation of insulin receptor-β and glucose transporter 4 (GLUT 4) translocation.



99.24% Purity:

Clinical Data: No Development Reported Size: 5 mg, 10 mg, 20 mg

## Ridaforolimus

(MK-8669; Deforolimus; AP23573)

Ridaforolimus (MK-8669) is a potent and selective mTOR inhibitor; inhibits ribosomal protein S6 phosphorylation with an IC<sub>so</sub> of 0.2 nM in HT-1080

Cat. No.: HY-50908

Purity: 97.83% Clinical Data: Phase 3 Size 10 mg, 50 mg

#### Rilmenidine

Purity:

Size:

Cat. No.: HY-100490

Rilmenidine, an innovative antihypertensive agent, is an orally active, selective I1 imidazoline receptor agonist. Rilmenidine is an alpha 2-adrenoceptor agonist. Rilmenidine induces autophagy.



#### Rilmenidine hemifumarate

Cat. No.: HY-100490A

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 induces autophagy.

Purity: 99.82% Clinical Data: Launched 5 mg, 10 mg Size

#### Rilmenidine phosphate

Clinical Data: Launched

>98%

1 mg, 5 mg

Cat. No.: HY-100490B

antihypertensive agent, is an orally active, selective I1 imidazoline receptor agonist. Rilmenidine phosphate is an alpha 2-adrenoceptor agonist. Rilmenidine phosphate induces autophagy.



Rilmenidine phosphate, an innovative

Purity: ≥98.0% Clinical Data: Launched Size 5 mg, 10 mg, 25 mg

#### Rilmenidine-d4

Rilmenidine-d4 is the deuterium labeled Rilmenidine. Rilmenidine, an innovative antihypertensive agent, is an orally active, selective I1 imidazoline receptor agonist. Rilmenidine is an alpha 2-adrenoceptor agonist. Rilmenidine induces autophagy.

Cat. No.: HY-100490S

Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg

#### Rimacalib

(SMP 114) Cat. No.: HY-100779

Rimacalib (SMP 114) is a Ca2+/calmodulin-dependent protein kinase II (CaMKII) inhibitor, with IC  $_{50}s$  of  ${\sim}1~\mu\text{M}$  for  $CaMKII\alpha$  to ~30  $\mu M$  for  $CaMKII\gamma$ .

Purity: 99 65% Clinical Data: Phase 2

Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

#### RITA

(NSC 652287) Cat. No.: HY-13424

RITA is an inhibitor of p53-HDM-2 interaction, binds to p53dN, with a K, of 1.5 nM, and also induces DNA-DNA cross-links.



Purity: 99 45%

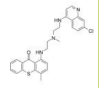
Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg

# **ROC-325**

Cat. No.: HY-103706

ROC-325 is a potent and orally active autophagy 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 apoptosis.



Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

#### Rosiglitazone

(BRL 49653) Cat. No.: HY-17386

Rosiglitazone (BRL 49653) is a selective, orally active PPARy agonist with EC<sub>50</sub>s of 30 nM, 100 nM and 60 nM for PPARy1, PPARy2, and PPARy, respectively. Rosiglitazone binds to PPARy with a

K<sub>d</sub> of approximately 40 nM.

**Purity:** 99.90% Clinical Data: Launched

10 mM × 1 mL, 50 mg, 200 mg



#### Rosiglitazone hydrochloride

(BRL 49653 hydrochloride) Cat. No.: HY-17386A

Rosiglitazone hydrochloride (BRL 49653 hydrochloride) is a selective, orally active PPARγ agonist with EC<sub>so</sub>s of 30 nM, 100 nM and 60 nM for PPARγ1, PPARγ2, and PPARγ, respectively. Rosiglitazone hydrochloride binds to PPARy with a K<sub>d</sub> of approximately 40 nM.



>98% Purity: Clinical Data: Launched Size: 1 mg, 5 mg

#### Rosiglitazone maleate

(BRL 49653C) Cat. No.: HY-14600

Rosiglitazone maleate (BRL 49653C) is a potent and selective activator of PPARy, with EC<sub>50</sub>s of 30 nM, 100 nM and 60 nM for PPARy1, PPARy2, and **PPARγ**, respectively, and a  $K_d$  of appr 40 nM for PPARy; Rosiglitazone maleate is also an modulator of TRP channels, inhibits TRP melastatin...

99.75% Purity: Clinical Data: Launched Size: 50 mg, 200 mg

#### Rosiglitazone-d3

Cat. No.: HY-17386S

Rosiglitazone-d3 (BRL 49653-d3) is the deuterium labeled Rosiglitazone. Rosiglitazone (BRL 49653) is a selective, orally active PPARy agonist with EC<sub>50</sub>s of 30 nM, 100 nM and 60 nM for PPARγ1, PPARγ2, and PPARγ, respectively.

Rosuvastatin (ZD 4522)

> Rosuvastatin (ZD 4522) is a competitive HMG-CoA reductase inhibitor with an IC<sub>50</sub> of 11 nM.



Cat. No.: HY-17504A

>98% Purity: Clinical Data: Launched Size 1 mg, 5 mg

>98% Purity:

Clinical Data:

Size: 1 mg, 5 mg

Rosuvastatin Calcium

(Rosuvastatin hemicalcium; ZD 4522 Calcium) Cat. No.: HY-17504

Rosuvastatin Calcium (Rosuvastatin hemicalcium) is a competitive HMG-CoA reductase inhibitor with an IC<sub>so</sub> of 11 nM.

Purity: 99.94% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg

#### Rosuvastatin D3

(ZD 4522 D3) Cat. No.: HY-17504AS

Rosuvastatin D3 (ZD 4522 D3) is a deuterium labeled Rosuvastatin. Rosuvastatin (ZD 4522) is a competitive HMG-CoA reductase inhibitor with an IC<sub>50</sub> of 11 nM.



Purity: >98%

Clinical Data: No Development Reported

1 mg

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#### Rosuvastatin D3 Sodium

Cat. No.: HY-17504BS

Rosuvastatin D3 Sodium is deuterium labeled Rosuvastatin, which is a competitive inhibitor of HMG-CoA reductase with IC50 of 11 nM.

**Purity:** >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

### Rosuvastatin D6 Sodium

Rosuvastatin D6 Sodium is deuterium labeled Rosuvastatin, which is a competitive inhibitor of HMG-CoA reductase with IC50 of 11 nM.



Cat. No.: HY-17504BS1

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Rosuvastatin-d6 calcium

Cat. No.: HY-17504S

Rosuvastatin D6 Calcium is deuterium labeled Rosuvastatin, which is a competitive inhibitor of HMG-CoA reductase with IC50 of 11 nM.

**Purity:** 98 54%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Rotenone

Cat. No.: HY-B1756

Rotenone is an mitochondrial electron transport chain complex I inhibitor. Rotenone induces apoptosis through enhancing mitochondrial reactive oxygen species production.



**Purity:** 99 65%

Clinical Data: No Development Reported

10 mM × 1 mL, 500 mg, 1 g, 5 g

#### Rottlerin

(Mallotoxin; NSC 56346; NSC 94525)

Rottlerin, a natural product purified from Mallotus Philippinensis, is a specific PKC inhibitor, with  $IC_{50}$  values for PKC $\delta$  of 3-6  $\mu M$ , PKCα,β, $\gamma$  of 30-42 μM, PKCε,η, $\zeta$  of 80-100 μM.

Cat. No.: HY-18980

Purity: 98.02%

Clinical Data: No Development Reported

Size: 10 mg, 25 mg

#### RSVA405

Cat. No.: HY-103238

RSVA405 is a potent, orally active activator of AMPK, with an EC<sub>50</sub> of 1  $\mu$ M. RSVA405 facilitates CaMKKβ-dependent activation of AMPK, inhibits mTOR, and promotes autophagy to increase Aß degradation.



99.56% Purity:

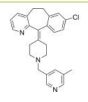
Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg Size:

#### Rupatadine

(UR-12592) Cat. No.: HY-13511

Rupatadine (UR-12592) is a potent, orally active and long-lasting dual PAF/H1 antagonist, with K,s of 0.55  $\mu$ M and 0.1  $\mu$ M, respectively. Rupatadine can be used for the research of allergic rhinitis and urticaria



>98% Purity: Clinical Data: Launched Size: 1 mg, 5 mg

## Rupatadine D4 fumarate

(UR-12592 D4 fumarate)

Cat. No.: HY-13511AS Rupatadine D4 fumarate (UR-12592 D4 fumarate) is a deuterium labeled Rupatadine fumarate. Rupatadine

Fumarate (UR-12592 Fumarate) is a potent dual PAF/H1 antagonist with K<sub>i</sub> of 0.55/0.1 μM(rabbit platelet membranes/quinea pig cerebellum membranes).

Purity: >98%

Clinical Data: No Development Reported

Size 1 mg, 5 mg



#### Rupatadine Fumarate

(UR-12592 Fumarate) Cat. No.: HY-13511A

Rupatadine (UR-12592) Fumarate is a potent, orally active and long-lasting dual PAF/H1 antagonist, with K<sub>i</sub>s of 0.55 μM and 0.1 μM, respectively. Rupatadine Fumarate can be used for the research of allergic rhinitis and urticaria.



Purity: 99.93% Clinical Data: Launched

Size 10 mM × 1 mL, 100 mg, 500 mg

#### Rutin

(Rutoside; Quercetin 3-O-rutinoside)

Rutin (Rutoside) is a flavonoid found in many plants and shows a wide range of biological activities including anti-inflammatory, antidiabetic, antioxidant, neuroprotective, nephroprotective, hepatoprotective and reducing  $\ensuremath{\mathsf{A}\beta}$ oligomer activities.



10 mM × 1 mL, 500 mg, 5 g, 10 g



Cat. No.: HY-N0148

#### Rutin hydrate

(Rutoside hydrate; Quercetin 3-O-rutinoside hydrate) Cat. No.: HY-N0148A

Rutin (Rutoside) hydrate is a flavonoid found in many plants and shows a wide range of biological activities including anti-inflammatory, antidiabetic, antioxidant, neuroprotective, nephroprotective, hepatoprotective and reducing A $\beta$  oligomer activities.

Purity: ≥98.0% Clinical Data: Launched Size: 500 mg



### Ruxolitinib phosphate

(INCB018424 phosphate)

Ruxolitinib phosphate (INCB018424 phosphate) is a potent <code>JAK1/2</code> inhibitor with <code>IC $_{50}$ </code> of 3.3 nM/2.8 nM, respectively, showing more than 130-fold selectivity over <code>JAK3</code>.



Cat. No.: HY-50858

Purity: 99.98% Clinical Data: Launched

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

# Ruxolitinib sulfate

(INCB018424 sulfate)

Clinical Data: Launched

Ruxolitinib

(INCB18424)

mitophagy.

Purity:

Ruxolitinib sulfate (INCB018424 sulfate) is the first potent, selective JAK1/2 inhibitor to enter the clinic with  $\rm IC_{50}$ S of 3.3 nM/2.8 nM, and has > 130-fold selectivity for JAK1/2 versus JAK3.

Ruxolitinib (INCB18424) is a potent and selective

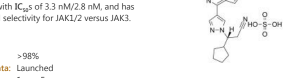
JAK1/2 inhibitor with IC<sub>sn</sub>s of 3.3 nM and 2.8 nM

in cell-free assays, and has 130-fold selectivity

autophagy and kills tumor cells through toxic

for JAK1/2 over JAK3. Ruxolitinib induces

99 99%



Purity: >98%
Clinical Data: Launched
Size: 1 mg, 5 mg

#### S130

Cat. No.: HY-112818

S130 is a high affinity, selective inhibitor of ATG4B (a major cysteine protease) with an  $IC_{50}$  of 3.24  $\mu$ M. S130 suppresses autophagy flux.

**Purity:** 99.31%

Clinical Data: No Development Reported

Size:  $10 \text{ mM} \times 1 \text{ mL}$ , 5 mg, 10 mg, 50 mg, 100 mg

## S29434

(NMDPEF) Cat. No.: HY-122614

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

S29434 (NMDPEF) is a potent, competitive, selective and cell-permeable inhibitor of **quinone reductase 2 (QR2)**, with  $\rm IC_{50}$ s ranging from 5 to 16 nM for human QR2 at different organizational levels, and has good selectivity for QR2 over QR1.



Cat. No.: HY-50856

Cat. No.: HY-50859

**Purity:** 99.37%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

#### SA 47

Cat. No.: HY-18080

SA 47 is a selective and potent inhibitor of fatty acid amide hydrolase (FAAH) and carbamate.

**Purity:** ≥99.0%

Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg

#### SA72

Cat. No.: HY-U00240

SA72 is a highly selective **fatty acid amide hydrolase** (FAAH) inhibitor.

**Purity:** >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Salbutamol hemisulfate

(Albuterol hemisulfate; AH-3365 hemisulfate) Cat. No.: HY-B0436

Salbutamol Hemisulfate (Albuterol hemisulfate) is a short-acting  $\beta 2$  adrenergic receptor agonist Target:  $\beta 2$  Adrenergic Receptor Salbutamol Hemisulfate (Albuterol hemisulfate) is a short-acting, selective beta2-adrenergic receptor agonist used in the treatment of asthma and...



Purity: ≥98.0% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg

#### Salicylic acid

(2-Hydroxybenzoic acid)

Salicylic acid (2-Hydroxybenzoic acid) inhibits cyclo-oxygenase-2 (COX-2) activity independently of transcription factor (NF-κB) activation.



Cat. No.: HY-B0167

Purity: 96.22% Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 10 g, 50 g

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#### Salicylic acid-d6

(2-Hydroxybenzoic acid-d6) Cat. No.

Salicylic acid-D6 (2-Hydroxybenzoic acid-D6) is a deuterium labeled Salicylic acid. Salicylic acid inhibits cyclo-oxygenase-2 (COX-2) activity independently of transcription factor (NF-κB) activation.

**Purity:** >98%

Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg

#### Cat. No.: HY-B0167S (Procoxacin)

Procoxacin) Cat. No.: HY-15597

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:** ≥98.0%

Salinomycin

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

# Salirasib (S-Farnesylthiosalicylic acid; Farnesyl

Thiosalicylic Acid; FTS)

Cat. No.: HY-14754

Salirasib is a Ras inhibitor that inhibits specifically both oncogenically activated Ras and growth factor receptor-mediated Ras activation, resulting in the inhibition of Ras-dependent tumor growth.



Purity: 99.01% Clinical Data: Phase 2

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg

#### Salinomycin sodium salt

(Salinomycin sodium; Sodium salinomycin) Cat. No.: HY-17439

Salinomycin sodium salt (Salinomycin sodium), an antibiotic potassium ionophore, is a potent inhibitor of  $Wnt/\beta$ -catenin signaling.



**Purity:** >98%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 25 mg, 50 mg, 100 mg

#### Salubrinal

Cat. No.: HY-15486

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:** 99.69%

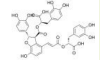
Clinical Data: No Development Reported

Size:  $10 \text{ mM} \times 1 \text{ mL}$ , 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

#### Salvianolic acid B

(Lithospermic acid B)

Salvianolic acid B is an active ingredient of Salvia miltiorrhiza, which has been widely applied in China for the management of various microcirculation-related disorders, such as cardiovascular disease, cerebrovascular disease, and diabetic vascular complication.



Cat. No.: HY-N1362

**Purity:** 99.73%

Clinical Data: No Development Reported
Size: 5 mg, 10 mg, 25 mg

#### Salvigenin

Cat. No.: HY-N1318

Salvigenin is a natural polyphenolic compound, with neuroprotective effect. Salvigenin has antitumor cytotoxic and immunomodulatory properties.

Purity: 99.79%

Clinical Data: No Development Reported

Size: 5 ma. 10 ma

#### Salvigenin-d9

Cat. No.: HY-N1318S
Salvigenin-d9 is the deuterium labeled Salvigenin.

Salvigenin is a natural polyphenolic compound, with neuroprotective effect. Salvigenin has antitumor cytotoxic and immunomodulatory properties.



**Purity:** >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Samotolisib

(LY3023414) Cat. No.: HY-12513

Samotolisib (LY3023414) potently and selectively inhibits class I PI3K isoforms, DNA-PK, and mTORC1/2 with IC $_{50}$ S of 6.07 nM, 77.6 nM, 38 nM, 23.8 nM, 4.24 nM and 165 nM for PI3K $\alpha$ , PI3K $\beta$ , PI3K $\beta$ , PI3K $\beta$ , PI3K $\beta$ , DNA-PK and mTOR, respectively.



Purity: 99.42% Clinical Data: Phase 2

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

#### Sanguinarine

(Sanguinarin; Sanguinarium; Pseudochelerythrine)

Sanguinarine (Sanguinarin), a benzophenanthridine alkaloid derived from the root of Sanguinaria Canadensis, can stimulate **apoptosis** via activating the production of reactive oxygen species (ROS). Sanguinarine-induced apoptosis is associated with the activation of JNK and NF-kB.



Cat. No.: HY-N0052

**Purity:** >98%

Clinical Data: No Development Reported
Size: 5 mg, 10 mg, 25 mg

#### Sanguinarine chloride (Sanguinarin chloride; Sanguinarium

chloride; Pseudochelerythrine chloride) Cat. No.: HY-N0052A

Sanguinarine (Sanguinarin) chloride, a benzophenanthridine alkaloid derived from the root of Sanguinaria Canadensis, can stimulate apoptosis via activating the production of reactive oxygen species (ROS). Sanguinarine-induced apoptosis is associated with the activation of JNK and NF-kB.



99 24% Purity:

Clinical Data: No Development Reported Size: 10 mM × 1 mL, 5 mg, 10 mg

Purity:

SAR405 is a first-in-class, selective, and ATP-competitive PI3K class III (PIK3C3) isoform Vps34 inhibitor ( $IC_{so}$ =1.2 nM;  $K_d$ =1.5 nM). SAR405 inhibits autophagy induced either by starvation or by mTOR inhibition. Anticancer activity.

**Purity:** 99.13%

Clinical Data: No Development Reported

10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 25 mg, 50 mg

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Saquinavir Mesylate

(Ro 31-8959/003) Cat. No.: HY-17003

Saguinavir mesylate is an HIV Protease Inhibitor used in antiretroviral therapy. IC50 Value: Target: HIV Protease Saquinavir is a protease inhibitor. Proteases are enzymes that cleave protein molecules into smaller fragments.



Purity: 99.89% Clinical Data: Launched

10 mM × 1 mL, 10 mg, 50 mg, 100 mg

#### Saracatinib

(AZD0530) Cat. No.: HY-10234

Saracatinib (AZD0530) is a potent Src family inhibitor with  $IC_{50}$ s of 2.7 to 11 nM for c-Src, Lck, c-YES, Lyn, Fyn, Fgr, and Blk. Saracatinib shows high selectivity over other tyrosine kinases.



99 97% Purity: Clinical Data: Phase 3

Size:  $10 \text{ mM} \times 1 \text{ mL}$ , 10 mg, 50 mg, 100 mg

#### SB 202190

SB 202190 is a selective p38 MAP kinase inhibitor with  $IC_{so}$ s of 50 nM and 100 nM for p38 $\alpha$  and p38 $\beta$ 2, respectively. SB 202190 binds to the ATP pocket of the active recombinant human p38 kinase with a K of 38 nM. SB 202190 has anti-cancer activity and rescued memory deficits.

99.89% **Purity:** 

SB 216763

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 50 mg, 100 mg, 200 mg

SB 216763 is potent, selective and ATP-competitive GSK-3 inhibitor with  $IC_{50}$ s of 34.3 nM for both

SB 202190 hydrochloride

Cat. No.: HY-10295A

SB 202190 hydrochloride is a selective p38 MAP kinase inhibitor with IC<sub>50</sub>s of 50 nM and 100 nM for p38α and p38β2, respectively. SB 202190 hydrochloride binds to the ATP pocket of the active recombinant human p38 kinase with a K<sub>d</sub> of 38 nM.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg Purity: 99.30%

GSK-3 $\alpha$  and GSK-3 $\beta$ .

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

SB 239063

Cat. No.: HY-11068

SB 239063 is a potent, selective and orally active p38 MAPK inhibitor, exhibits an IC<sub>so</sub> of 44 nM for recombinant purified human p38α, with equipotent inhibitory activity against  $p38\alpha$  and p38 $\beta$ . SB 239063 has no effect on p38 $\gamma$  or p38 $\delta$ .



99.80% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 25 mg, 50 mg

#### SB 242235

SB-242235 is a potent and selective p38 MAP kinase inhibitor, with an  $IC_{50}$  of 1.0 $\mu M$  in primary human chondrocytes.

99.51%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg



Cat. No.: HY-13328

**SAR405** 

Sapanisertib

(INK-128; MLN0128; TAK-228)

Sapanisertib (INK-128; MLN0128; TAK-228) is an

inhibitor with an IC<sub>so</sub> of 1 nM for mTOR kinase.

orally available, ATP-dependent mTOR1/2

99 66%

Clinical Data: Phase 2

Cat. No.: HY-12481

Cat. No.: HY-10295



Cat. No.: HY-12012

Cat. No.: HY-18306

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#### SBE13

Cat. No.: HY-15158A

SBE13 is a potent and selective Plk1 inhibitor, with an IC<sub>so</sub> of 200 pM; SBE13 poorly inhibits Plk2  $(IC_{50}>66 \mu M)$  or Plk3  $(IC_{50}=875 \text{ nM})$ .

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

## SBE13 Hydrochloride

SBE13 Hydrochloride is a potent and selective Plk1

inhibitor, with an IC<sub>50</sub> of 200 pM; SBE13 Hydrochloride poorly inhibits Plk2  $(IC_{50} > 66 \mu M)$  or Plk3  $(IC_{50} = 875 nM)$ .



Cat. No.: HY-15158

98 76% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 10 mg, 50 mg

#### SBI-0206965

Cat. No.: HY-16966

SBI-0206965 is a potent, selective and cell permeable autophagy kinase ULK1 inhibitor with IC<sub>so</sub>s of 108 nM for ULK1 kinase and 711 nM for the highly related kinase ULK2.



Purity: 99 39%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

#### SBI-0640756

(SBI-756) Cat. No.: HY-19560

SBI-0640756 (SBI-756) is an inhibitor of eIF4G1 and disrupts the eIF4F complex.



**Purity:** 99 76%

Clinical Data: No Development Reported

10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 25 mg, 50 mg

#### SBP-7455

Cat. No.: HY-137742

SBP-7455 is a potent, high affinity and orally active dual ULK1/ULK2 autophagy inhibitor with IC<sub>so</sub>s of 13 nM and 476 nM in the ADP-Glo assays, respectively. SBP-7455 potently inhibits ULK1/2 enzymatic activity and can be used for triple-negative breast cancer (TNBC) research.



98.29% Purity:

Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

#### Schaftoside

Schaftoside is a flavonoid found in a variety of Chinese herbal medicines, such as Eleusine indica. Schaftoside inhibits the expression of TLR4 and Myd88. Schaftoside also decreases Drp1 expression and phosphorylation, and reduces mitochondrial

fission.

**Purity:** 99.88%

Clinical Data: No Development Reported Size 10 mM × 1 mL, 5 mg, 10 mg, 20 mg



Cat. No.: HY-N0703

#### Schisandrin

#### (Schizandrin; Schizandrol; Schizandrol-A) Cat. No.: HY-N0691

Schisandrin (Schizandrin), a dibenzocyclooctadiene lignan, is isolated from the fruit of Schisandra chinensis Baill. Schisandrin exhibits antioxidant, hepatoprotective, anti-cancer and anti-inflammatory activities. Schisandrin also can reverses memory impairment in rats.



Purity: 99.51%

Clinical Data: No Development Reported 5 mg, 10 mg, 50 mg, 100 mg Size:

#### Schisandrin A

(Schizandrin-A; Wuweizisu-A; Deoxyschizandrin) Cat. No.: HY-N0693

Schisandrin A inhibits CYP3A activity with an  $IC_{so}$  of 6.60  $\mu$ M and  $K_i$  of 5.83  $\mu$ M, respectively.



99.43% Purity:

Clinical Data: No Development Reported 10 mM × 1 mL, 10 mg Size:

#### Schisandrin B

#### (γ-Schisandrin; Wuweizisu B) Cat. No.: HY-N0089

Schisandrin B (y-Schisandrin) is a dibenzocyclooctadiene derivative isolated from Fructus Schisandrae, has been shown to produce antioxidant effect on rodent liver and heart.



Purity: 99.86%

No Development Reported Clinical Data: Size: 10 mM × 1 mL, 10 mg, 50 mg

#### Schisandrol B

#### (Gomisin-A; TJN-101; Wuweizi alcohol-B)

Schisandrol B (Gomisin-A) is a major active constituent of Schisandra sphenanthera with hepato-protective effects. Schisandrol B inhibits reactive oxygen species (ROS) production.



Cat. No.: HY-N0692

99.57%

Clinical Data: No Development Reported 10 mM × 1 mL, 10 mg

#### Scriptaid

(Scriptide; GCK1026) Cat. No.: HY-15489

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

#### Scutellarein

(6-Hydroxyapigenin; 4',5,6,7-Tetrahydroxyflavone)

Scutellarin, a main active ingredient extracted from Erigeron breviscapus (Vant.) Hand-Mazz., has been wildly used to treat acute cerebral infarction and paralysis induced by cerebrovascular diseases.



Cat. No.: HY-N0752

99 75% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 25 mg, 50 mg

#### SD 0006

(SD-06) Cat. No.: HY-11087

SD 0006 (SD-06) is an orally active, selective, ATP-competitive and potent diaryl pyrazole inhibitor of p38 $\alpha$  MAP kinase, with an IC<sub>50</sub> of 110 nM for p38 $\alpha$ .

Purity: 98 60%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

#### Sedanolide

Sedanolide, a natural compound occurring in edible umbelliferous plants, possesses anti-inflammatory and antioxidant activities.



Cat. No.: HY-N2114

**Purity:** >98%

Clinical Data: No Development Reported 5 mg, 10 mg, 25 mg

#### Selenomethionine

(Seleno-DL-methionine; DL-Selenomethionine) Cat. No.: HY-B1000

Selenomethionine is a naturally occurring amino acid containing selenium, is a common natural food source of selenium.

Purity: > 98.0% Clinical Data: Launched

10 mM × 1 mL, 100 mg, 500 mg Size:

#### Sepantronium bromide

(YM-155) Cat. No.: HY-10194

Sepantronium bromide (YM-155) is a survivin inhibitor with an IC<sub>50</sub> of 0.54 nM.



98.91% Purity: Clinical Data: Phase 2

Size  $10~\text{mM}\times1~\text{mL},\,5~\text{mg},\,10~\text{mg},\,50~\text{mg},\,100~\text{mg}$ 

#### Sepantronium hydrochloride

(YM-155 hydrochloride) Cat. No.: HY-10194A

Sepantronium hydrochloride (YM-155 hydrochloride) is a novel survivin suppressant with an IC<sub>so</sub> of 0.54 nM for the inhibition of survivin promoter activity.

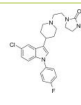
>98% Purity: Clinical Data: Phase 2 Size: 1 mg, 5 mg

#### Sertindole (Lu 23-174)

Sertindole, a neuroleptic, is one of the newer antipsychotic medications available. Target: Multi-target In vitro studies showed that sertindole exerts a potent antagonism at serotonin 5-HT2A, 5-HT2C, dopamine D2, and αl adrenergic receptors.

Purity: 99.76% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg



Cat. No.: HY-14543

#### Sertindole-d4

Cat. No.: HY-14543S

Sertindole-d4 (Lu 23-174-d4) is the deuterium labeled Sertindole. Sertindole, a neuroleptic, is one of the newer antipsychotic medications available.



Purity: >98% Clinical Data: Size: 1 mg

#### Sevelamer

Sevelamer is a phosphate binding drug used to treat hyperphosphatemia in patients with chronic kidney disease; consists of polyallylamine that is crosslinked with epichlorohydrin.

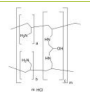
Cat. No.: HY-13995

Purity: >98% Clinical Data: Launched 1 mg, 5 mg

#### Sevelamer hydrochloride

Sevelamer hydrochloride is a phosphate binding drug used to treat hyperphosphatemia in patients with chronic kidney disease; consists of polyallylamine that is crosslinked with epichlorohydrin.

Purity: >98.0% Clinical Data: Launched 100 mg, 500 mg Size:

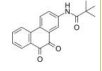


Cat. No.: HY-13995A

# SF1670

Cat. No.: HY-15842

SF1670 is a potent and specific phosphatase and tensin homolog deleted on chromosome 10 (PTEN) inhibitor.



Purity: > 98.0%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg Size

#### SH379

Cat. No.: HY-143656

SH379 is the derivative of 2-methylpyrimidine-fused tricyclic diterpene. SH379 is a potent and orally active anti-late-onset hypogonadism agent.



>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Shogaol

([6]-Shogaol; 6-Shogaol) Cat. No.: HY-14616

Shogaol (-Shogaol), an active compound isolated from Ginger (Zingiber officinale Rosc), exhibits a variety of biological activities including anticancer, anti-inflammation, and anti-oxidation.



99.84% Purity:

Clinical Data: No Development Reported 10 mM  $\times$  1 mL, 5 mg, 10 mg, 25 mg Size

## Sildenafil citrate

(UK-92480 citrate) Cat. No.: HY-15025A

Sildenafil citrate is a potent phosphodiesterase type 5 (PDE5) inhibitor with  $IC_{50}$  of 5.22 nM.



Purity: 99.73% Launched Clinical Data:

Size: 10 mM × 1 mL, 50 mg, 100 mg, 200 mg, 500 mg

#### Sevelamer-(d5)n hydrochloride

Sevelamer-(d5)n hydrochloride is the deuterium labeled Sevelamer hydrochloride. Sevelamer hydrochloride is a phosphate binding drug used to treat hyperphosphatemia in patients with chronic kidney disease; consists of polyallylamine that is crosslinked with epichlorohydrin.

Cat. No.: HY-13995AS

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### SGI-1776

Cat. No.: HY-13287

SGI-1776 is an inhibitor of Pim kinases, with IC<sub>so</sub>s of 7 nM, 363 nM, and 69 nM for Pim-1, -2 and -3, respectively.



Purity: 99.23% Clinical Data: Phase 1

10 mM × 1 mL, 5 mg, 10 mg, 50 mg

#### SH498

Cat. No.: HY-143658

SH498, a novel Bmi-1-mediated antitumor agent, shows potent antiproliferative activity.



>98% Purity:

Clinical Data: No Development Reported

Size 1 mg, 5 mg

#### Sildenafil

(UK-92480) Cat. No.: HY-15025

Sildenafil (UK-92480) is a potent phosphodiesterase type 5 (PDE5) inhibitor with an IC<sub>so</sub> of 5.22 nM.



99.90% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 50 mg, 100 mg, 200 mg

#### Sildenafil-d8

(UK-92480-d8) Cat. No.: HY-15025S1

Sildenafil-d8 (UK-92480-d8) is the deuterium labeled Sildenafil. Sildenafil (UK-92480) is a potent phosphodiesterase type 5 (PDE5) inhibitor with an IC<sub>50</sub> of 5.22 nM.



**Purity:** >98%

Clinical Data: No Development Reported

1 mg, 10 mg

#### Silibinin

(Silibinin A; Silymarin I) Cat. No.: HY-13748

Silibinin (Silibinin A), an effective anti-cancer and chemopreventive agent, has been shown to exert multiple effects on cancer cells, including inhibition of both cell proliferation and migration.



Purity: 99.87% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg

#### Silmitasertib

(CX-4945) Cat. No.: HY-50855

Silmitasertib (CX-4945) is an orally bioavailable, highly selective and potent CK2 inhibitor, with  $IC_{s0}$  values of 1 nM against CK2 $\alpha$  and CK2 $\alpha$ '.



Purity: 99.92% Clinical Data: Phase 2

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

#### Silmitasertib sodium salt

(CX-4945 sodium salt) Cat. No.: HY-50855B

Silmitasertib sodium salt is an orally bioavailable, highly selective and potent CK2 inhibitor, with  $IC_{50}$  values of 1 nM against CK2 $\alpha$  and CK2 $\alpha$ '.

Purity: 99.93% Clinical Data: Phase 2

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg

#### Silvestrol

((-)-Silvestrol) Cat. No.: HY-13251

Silvestrol is a eukaryotic translation initiation factor 4A (eIF4A) inhibitor isolated from the fruits and twigs of Aglaia foveolata. Silvestrol induces autophagy and caspase-mediated apoptosis.



Purity: 98.11%

Clinical Data: No Development Reported Size: 1 mg, 2 mg, 5 mg, 10 mg

#### Simvastatin

(MK 733) Cat. No.: HY-17502

Simvastatin (MK 733) is a competitive inhibitor of HMG-CoA reductase with a  $K_i$  of 0.2 nM.



Purity: 99.45% Clinical Data: Launched

Size: 50 mg, 100 mg, 200 mg, 500 mg

## Simvastatin-d6

(MK 733-d6) Cat. No.: HY-110231

Simvastatin-d6 (MK 733-d6) is the deuterium labeled Simvastatin. Simvastatin (MK 733) is a competitive inhibitor of **HMG-CoA reductase** with a **K**, of 0.2 nM.



**Purity:** >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Sinomenine

Cat. No.: HY-15122

Sinomenine, an alkaloid extracted from Sinomenium acutum, is a blocker of the  $NF-\kappa B$  activation. Sinomenine also is an activator of  $\mu$ -opioid receptor.



Purity: 99.88% Clinical Data: Launched

Size:  $10 \text{ mM} \times 1 \text{ mL}, 100 \text{ mg}$ 

#### Sinomenine hydrochloride

(Cucoline hydrochloride) Cat. No.: HY-15122A

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.



Purity: 99.88%

Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg

# OH HCI

#### Sirtinol

Cat. No.: HY-13515

Sirtinol is a **sirtuin** (SIRT) inhibitor, with IC $_{50}$ s of 48  $\mu$ M, 57.7  $\mu$ M and 131  $\mu$ M for ySir2, hSIRT2 and hSIRT2, respectively.



**Purity:** ≥98.0%

Clinical Data: No Development Reported

Size:  $10 \text{ mM} \times 1 \text{ mL}$ , 5 mg, 10 mg, 50 mg, 100 mg

#### Sitagliptin (MK-0431)

Sitagliptin (MK-0431) is a potent inhibitor of

DPP4 with an  $IC_{50}$  of 19 nM in Caco-2 cell extracts.



Cat. No.: HY-13749

Purity: 99.72% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 200 mg

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#### Sitagliptin phosphate

(MK-0431 phosphate) Cat. No.: HY-13749A

Sitagliptin phosphate (MK-0431 phosphate) is a potent inhibitor of DPP4 with an IC<sub>so</sub> of 19 nM in Caco-2 cell extracts.

Purity: >99.0% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 200 mg

## Sitagliptin phosphate monohydrate

(MK-0431 phosphate monohydrate)

Sitagliptin phosphate monohydrate (MK-0431 phosphate monohydrate) is a potent inhibitor of DPP4 with an IC<sub>50</sub> of 19 nM in Caco-2 cell



Cat. No.: HY-13749B

Purity: 99 62% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 200 mg

## Sitagliptin-d4 phosphate

Cat. No.: HY-13749AS

Sitagliptin-d4 (MK-0431-d4) phosphate is the deuterium labeled Sitagliptin phosphate. Sitagliptin phosphate (MK-0431 phosphate) is a potent inhibitor of **DPP4** with an **IC**<sub>50</sub> of 19 nM in Caco-2 cell extracts.



Purity: >98%

Clinical Data: Size: 1 mg

# $SJF\alpha$

SJFα is a 13-atom linker **PROTAC** based on **von** Hippel-Lindau ligand. SJF $\alpha$  degrades p38 $\alpha$  with a DC<sub>so</sub> of 7.16nM, but is far less effective at degrading p38 $\delta$  (DC<sub>50</sub>=299nM) and does not degrade the other p38 isoforms (β and γ) at

concentrations up to 2.5µM. >98% **Purity:** 

Clinical Data: No Development Reported

1 mg, 5 mg

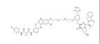


Cat. No.: HY-114404

SJFδ

Cat. No.: HY-114405

SJF $\delta$  is a 10-atom linker **PROTAC** based on **von** Hippel-Lindau ligand. SJFδ degrades p38δ with a  $DC_{so}$  of 46.17nM, but does not degrade p38 $\alpha$ , p38β, or p38γ.



Purity: >98%

Clinical Data: No Development Reported

Size: 5 mg

#### Skatole

(3-Methylindole; 3-Methyl-1H-indole)

Skatole is produced by intestinal bacteria, regulates intestinal epithelial cellular functions through activating aryl hydrocarbon receptors and p38.



Cat. No.: HY-W007355

99.86% Purity:

Clinical Data: No Development Reported Size 10 mM × 1 mL, 100 mg

#### Skatole-d3

#### (3-Methylindole-d3; 3-Methyl-1H-indole-d3) Cat. No.: HY-W007355S

Skatole-d3 (3-Methylindole-d3) is the deuterium labeled Skatole. Skatole is produced by intestinal bacteria, regulates intestinal epithelial cellular functions through activating aryl hydrocarbon receptors and p38.



>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Skatole-d8

(3-Methylindole-d8; 3-Methyl-1H-indole-d8)

Skatole-d8 (3-Methylindole-d8) is the deuterium labeled Skatole. Skatole is produced by intestinal bacteria, regulates intestinal epithelial cellular functions through activating aryl hydrocarbon receptors and p38.



Cat. No.: HY-W007355S1

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

# SKF-96365 hydrochloride

Cat. No.: HY-100001

SKF-96365 hydrochloride is a potent TRP channel blocker and a store-operated Ca<sup>2+</sup> entry (SOCE) 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.



Purity: 99.51%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

#### Skepinone-L

(CBS3830) Cat. No.: HY-15300

Skepinone-L (CBS3830) is a selective p38 mitogen-activated protein kinase inhibitor.



Purity: 99.77%

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

#### SLLN-15

SLLN-15 is an oral active, selective and potent enhancer of autophagy that activates cytostatic macroautophagy/autophagy in triple-negative breast

cancer (TNBC).

Cat. No.: HY-125465

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

## SMER18

SMER18 is a small molecule enhancer of rapamycin which act as a mTOR-independent autophagy inducer.



Cat. No.: HY-18672

98 60% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

# SMER28

Cat. No.: HY-100200

SMER28 is a positive regulator of autophagy acting via an mTOR-independent mechanism. SMER28 prevents the accumulation of amyloid beta peptide.



Purity: 99 99%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg, 200 mg

#### SN-38

(NK012) Cat. No.: HY-13704

SN-38 (NK012) is an active metabolite of the Topoisomerase I inhibitor Irinotecan. SN-38 (NK012) inhibits DNA and RNA synthesis with IC<sub>50</sub>s of 0.077 and 1.3 µM, respectively.



**Purity:** 99 80% Clinical Data: Phase 2

10 mM × 1 mL, 50 mg, 100 mg, 200 mg, 500 mg

## SN-38-d3

(NK012-d3) Cat. No.: HY-13704S

SN-38-d3 is the deuterium labeled SN-38. SN-38 (NK012) is an active metabolite of the Topoisomerase I inhibitor Irinotecan. SN-38 (NK012) inhibits DNA and RNA synthesis with ICsos of 0.077 and 1.3 µM, respectively.



>98% Purity:

Clinical Data: No Development Reported

Size: 1 ma

# SN-38-d5

(NK012-d5) Cat. No.: HY-13704S1

SN-38-d5 is deuterium labeled SN-38. SN-38 (NK012) is an active metabolite of the Topoisomerase I inhibitor Irinotecan. SN-38 (NK012) inhibits DNA and RNA synthesis with IC50s of 0.077 and 1.3  $\mu M$ , respectively.



>98% Purity:

Clinical Data: No Development Reported

Size 1 mg, 5 mg

Sodium 4-phenylbutyrate (4-PBA sodium; 4-Phenylbutyric acid sodium; Benzenebutyric acid sodium) Cat. No.: HY-15654

Sodium 4-phenylbutyrate (4-PBA sodium) is an

inhibitor of HDAC and endoplasmic reticulum (ER) stress, used in cancer and infection research.

99.96% Purity: Clinical Data: Launched 100 mg, 200 mg Size

#### Sodium Salicylate (Salicylic acid sodium salt;

2-Hydroxybenzoic acid sodium salt)

Sodium Salicylate (Salicylic acid sodium salt) inhibits cyclo-oxygenase-2 (COX-2) activity independently of transcription factor (NF-κB) activation. Sodium Salicylate is also a S6K inhibitor



Cat. No.: HY-B0167A

Purity: 99.88% Clinical Data: Launched

10 mM × 1 mL, 500 mg, 10 g, 50 g

#### Sofalcone

Cat. No.: HY-B2184

Sofalcone, a gastric antiulcer agent, is known to induce the expression of Heme oxygenase-1 (HO-1) in gastric epithelium.

Purity: 99.12% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg, 1 g, 5 g

#### Sophocarpine

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.

Purity: ≥98.0%

Clinical Data: No Development Reported 10 mM × 1 mL, 5 mg, 10 mg, 20 mg



Cat. No.: HY-N0103

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#### Sophocarpine monohydrate

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.



Cat. No.: HY-N0103A

H<sub>2</sub>O

Purity: 99.15%

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 5 mg, 10 mg

# Sorafenib Tosylate

(Bay 43-9006 Tosylate) Cat. No.: HY-10201A

Sorafenib Tosylate (Bay 43-9006 Tosylate) is a potent and orally active Raf inhibitor with  $IC_{50}$ s of 6 nM and 20 nM for Raf-1 and B-Raf, respectively.



Purity: 99.75%
Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg

Sorafenib-d3

(Bay 43-9006-d3; Donafenib) Cat. No.: HY-10201S

Sorafenib-d3 (Bay 43-9006-d3) 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.

Purity: 99.57% Clinical Data: Launched

Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

#### Soyasapogenol B

Soyasapogenol B, an ingredient of soybean, exerts anti-proliferative, anti-metastatic activities. Soyasapogenol B triggers endoplasmic reticulum stress, which mediates apoptosis and autophagy in

colorectal cancer.

**Purity:** 98.52%

Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg

# H OH

Cat. No.: HY-N6074

## Spautin-1

Cat. No.: HY-12990

Spautin-1 is a specific and potent **autophagy** inhibitor which inhibits ubiquitin-specific peptidases, USP10 and USP13 with  $IC_{50}$ s of 0.6-0.7  $\mu$ M.

**Purity:** 99.26%

Clinical Data: No Development Reported

Size:  $10 \text{ mM} \times 1 \text{ mL}$ , 5 mg, 10 mg, 50 mg, 100 mg

#### Sorafenib

(Bay 43-9006) Cat. No.: HY-10201

Sorafenib (Bay 43-9006) is a potent and orally active Raf inhibitor with  $\rm IC_{50}S$  of 6 nM and 20 nM for Raf-1 and B-Raf, respectively. Sorafenib is a multikinase inhibitor with  $\rm IC_{50}S$  of 90 nM, 15 nM, 20 nM, 57 nM and 58 nM for VEGFR2, VEGFR3, PDGFRB, FLT3 and c-Kit, respectively.

Purity: 99.92% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg

#### Sorafenib-13C,d3

Cat. No.: HY-10201S2

Sorafenib-13C,d3 is the 13C- and deuterium labeled Sorafenib. Sorafenib (Bay 43-9006) is a potent and orally active Raf inhibitor with IC $_{50}$ S of 6 nM and 20 nM for Raf-1 and B-Raf, respectively.



Purity: >98%

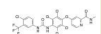
Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Sorafenib-d4

(Bay 43-9006-d4) Cat. No.: HY-10201S1

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.



**Purity:** >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### SP600125

Cat. No.: HY-12041

0

SP600125 is an orally active, reversible, and ATP-competitive JNK inhibitor with IC $_{\rm 50}$ s of 40, 40 and 90 nM for JNK1, JNK2 and JNK3, respectively. SP600125 is a potent ferroptosis inhibitor. SP600125 inhibits autophagy and activates apoptosis.

**Purity:** 99.55%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg, 500 mg

# Spironolactone

(SC9420) Cat. No.: HY-B0561

Spironolactone (SC9420) is an orally active aldosterone mineralocorticoid receptor antagonist with an IC $_{50}$  of 24 nM. Spironolactone is also a potent antagonist of androgen receptor with an IC $_{50}$  of 77 nM. Spironolactone promotes autophagy in podocytes.

Purity: 99.75% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 1 g, 5 g



#### Spironolactone-d3

(SC9420-d3) Cat. No.: HY-B0561S1

Spironolactone-d3 (SC9420-d3) is the deuterium labeled Spironolactone. Spironolactone (SC9420) is an orally active aldosterone mineralocorticoid receptor antagonist with an IC<sub>50</sub> of 24 nM.



>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Spironolactone-d7

(SC9420-d7) Cat. No.: HY-B0561S

Spironolactone-d7 (SC9420-d7) is the deuterium labeled Spironolactone. Spironolactone (SC9420) is an orally active aldosterone mineralocorticoid receptor antagonist with an IC<sub>50</sub> of 24 nM.



Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg

Purity:

SR-18292

Size:

SR-18292 is a PPAR gamma coactivator-1α (**PGC-1α**) inhibitor, which increases PGC-1α acetylation, suppresses gluconeogenic gene expression and reduces glucose production in hepatocytes.



Cat. No.: HY-101491

Cat. No.: HY-B0561S2

**Purity:** ≥98.0%

Spironolactone-d3-1 (SC9420-d3-1)

Spironolactone-d3-1 is deuterium labeled

orally active aldosterone mineralocorticoid

receptor antagonist with an IC50 of 24 nM.

Clinical Data: No Development Reported

1 mg, 5 mg

>98%

Spironolactone. Spironolactone (SC9420) is an

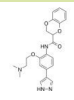
Clinical Data: No Development Reported

5 mg, 10 mg, 25 mg, 50 mg, 100 mg Size:

#### SR-3677

Cat. No.: HY-13300

SR-3677 is a potent and selective ROCK-II inhibitor with an  $IC_{50}$  of ~3 nM.



99.90% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 10 mg, 50 mg, 100 mg Size:

#### SR12813

(GW 485801) Cat. No.: HY-100793

SR12813 (GW 485801) is an inhibitor of 3-hydroxy-3-methylglutaryl-coenzyme A (HMG-CoA) reductase, with an  $IC_{50}$  value of 0.85  $\mu$ M. SR12813 is also an efficient agonist of human pregnane X receptor (hPXR). SR12813 can strongly bind to



99.39% Purity:

Clinical Data: No Development Reported

hPXR but not to mouse PXR (mPXR).

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

#### SRT 1720

Cat. No.: HY-10532

SRT 1720 is a selective activator of human SIRT1 with an  $EC_{_{1.5}}$  of 0.16  $\mu M\text{,}$  and shows less potent activities agaiinst SIRT2 and SIRT3 with EC<sub>1.5</sub>s of 37  $\mu$ M and > 300  $\mu$ M, respectively.



99.82% Purity:

Clinical Data: No Development Reported 10 mM  $\times$  1 mL, 10 mg, 50 mg Size:

#### SRT 1720 Hydrochloride

Cat. No.: HY-15145

SRT 1720 Hydrochloride is a selective activator of SIRT1 with an EC<sub>50</sub> of 0.10  $\mu$ M, and shows less potent activities on SIRT2 and SIRT3.



99.92% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg Size:

#### Stauprimide

Cat. No.: HY-N6747

Stauprimide is a staurosporine analog that promotes embryonic stem cell (ESC) differentiation.



Purity: ≥98.0%

Clinical Data: No Development Reported

Size: 100 μg, 500 μg

## Stavudine

(d4T) Cat. No.: HY-B0116

Stavudine (d4T) is an orally active nucleoside reverse transcriptase inhibitor (NRTI). Stavudine has activity against HIV-1 and HIV-2. Stavudine also inhibits the replication of mitochondrial DNA (mtDNA).



99.67% Purity: Clinical Data: Launched

10 mM × 1 mL, 100 mg, 500 mg

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#### Stavudine sodium

(d4T sodium) Cat. No.: HY-B0116A

Stavudine (d4T) sodium is an orally active nucleoside reverse transcriptase inhibitor (NRTI). Stavudine sodium has activity against HIV-1 and HIV-2. Stavudine sodium also inhibits the replication of mitochondrial DNA (mtDNA).

HN Na<sup>+</sup>

Purity: >98%
Clinical Data: Launched
Size: 1 mg, 5 mg

## Stavudine-d4

Stavudine-d4 is the deuterium labeled Stavudine. Stavudine (d4T) is an orally active **nucleoside reverse transcriptase** inhibitor (NRTI). Stavudine has activity against **HIV-1** and **HIV-2**. Stavudine also inhibits the replication of mitochondrial DNA (mtDNA).

O NO OF

Cat. No.: HY-B0116S

**Purity:** >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### STF-31

Cat. No.: HY-18728

STF-31 is a selective inhibitor of glucose transporter 1 (GLUT1), with an  $IC_{50}$  of  $1\mu M$ .

**Purity:** 96.97%

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 10 mg, 50 mg

#### STF-62247

Cat. No.: HY-100746

STF-62247 is an <code>autophagy</code> inducer that selectively cytotoxic to VHL-deficient renal cell carcinoma (IC $_{50}$  of 0.625  $\mu$ M and 16  $\mu$ M in RCC4 and RCC4/VHL cells, respectively).

N S H

Purity: 97.20%

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

#### Size: 10 m

STO-609

Cat. No.: HY-19805

STO-609 is a selective and cell-permeable inhibitor of the  $\text{Ca}^{2+}/\text{calmodulin-dependent}$  protein kinase kinase (CaM-KK), with K<sub>1</sub> values of 80 and 15 ng/mL for recombinant CaM-KK $\alpha$  and CaM-KK $\beta$ , respectively.

OH OH

Purity: 98.13%

Clinical Data: No Development Reported
Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

#### Streptozocin

(Streptozotocin; U 9889)

Streptozocin is a potent DNA-methylating antibiotic. Streptozotocin causes methylation of liver and kidney and pancreatic DNA, but no methylation in brain DNA.

HO OH OH OH O

Cat. No.: HY-13753

Purity: 99.15% Clinical Data: Launched Size: 100 mg, 500 mg

#### SU11274

(PKI-SU11274) Cat. No.: HY-12014

SU11274 is a selective Met inhibitor with  ${\rm IC}_{\rm 50}$  of 10 nM, but has no effects on PGDFR $\beta$ , EGFR or Tie2.

**Purity:** 98.19%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

#### SU9516

Cat. No.: HY-18629

SU9516 is a potent CDK2 inhibitor, with an  $\rm IC_{50}$  of 22 nM, and also shows inhibitory effects on CDK1 and CDK4, with  $\rm IC_{50}s$  of 40, 200 nM, respectively.



**Purity:** 99.83%

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

#### Sulfabenzamide

(N-Sulfanilylbenzamide) Cat. No.: HY-B0960

Sulfabenzamide (N-Sulfanilylbenzamide) is an antimicrobial agent and usually consumed in combination with Sulfathiazole and Sulfacetamide. Sulfabenzamide is effective against Gram-positive and negative bacterial strains.



Purity: 99.55% Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg

#### Sulfasalazine

(NSC 667219)

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.



Cat. No.: HY-14655

Purity: 99.04% Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g

#### Sulfasalazine-d4

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-κB activity. Sulfasalazine is a type 1 ferroptosis inducer.

Cat. No.: HY-14655S

Purity: >98%

Clinical Data: No Development Reported

Size: 2.5 mg, 25 mg

#### Sulindac-d3

(MK-231-d3) Cat. No.: HY-B0008S

Sulindac-d3 is deuterium labeled Sulindac. Sulindac (MK-231) is a non-steroidal antiinflammatory agent, acts as a COX-2 inhibitor, and inhibits overexpression of COX-2.

**Purity:** >98%

Clinical Data: No Development Reported

1 mg, 5 mg

# Sunitinib

Purity:

Size:

Sulindac

(MK-231)

Sulindac (MK-231) is a non-steroidal

99 81%

Clinical Data: Launched

antiinflammatory agent, acts as a COX-2

inhibitor, and inhibits overexpression of COX-2.

(SU 11248) Cat. No.: HY-10255A

Sunitinib (SU 11248) is a multi-targeted receptor tyrosine kinase inhibitor with IC<sub>50</sub>s of 80 nM and 2 nM for VEGFR2 and PDGFRβ, respectively.

10 mM × 1 mL, 100 mg, 500 mg



Cat. No.: HY-B0008

**Purity:** 98 96% Clinical Data: Launched

10 mM × 1 mL, 100 mg, 200 mg, 500 mg

#### Sunitinib Malate

(SU 11248 Malate) Cat. No.: HY-10255

Sunitinib Malate (SU 11248 Malate) is a multi-targeted receptor tyrosine kinase inhibitor with IC<sub>so</sub>s of 80 nM and 2 nM for VEGFR2 and PDGFRβ, respectively.



99 47% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 200 mg, 500 mg

# Sunitinib-d10

(SU 11248-d10) Cat. No.: HY-10255AS

Sunitinib D10 (SU 11248 D10) is a deuterium labeled Sunitinib. Sunitinib is a multi-targeted receptor tyrosine kinase inhibitor with IC so of 80 nM and 2 nM for VEGFR2 and PDGFRB, respectively.



99.89% Purity:

Clinical Data: No Development Reported

Size 1 mg, 5 mg

#### Sunitinib-d4

Cat. No.: HY-10255AS1

Sunitinib-d4 (SU 11248-d4) is the deuterium labeled Sunitinib. Sunitinib (SU 11248) is a multi-targeted receptor tyrosine kinase inhibitor with IC<sub>so</sub>s of 80 nM and 2 nM for VEGFR2 and PDGFRβ, respectively.



>98% Purity:

Clinical Data:

Size: 2.5 mg, 1 mg, 25 mg

#### Syntide 2

Cat. No.: HY-P0271

Syntide 2, a Ca2+- and calmodulin (CaM)-dependent protein kinase II (CaMKII) substrate peptide, selectively inhibits the gibberellin (GA) response, leaving constitutive and abscisic acid-regulated events unaffected.

PLARTLSVAGLPGKK

>98% Purity:

Clinical Data: No Development Reported 1 mg, 5 mg, 10 mg Size:

#### Syringin

(Eleutheroside B) Cat. No.: HY-N0824

Syringin is a main bioactive phenolic glycoside in Acanthopanax senticosus, with anti-osteoporosis activity. Syringin prevents cardiac hypertrophy induced by pressure overload through the attenuation of autophagy.



Purity: 99.05%

Clinical Data: No Development Reported Size: 5 mg, 10 mg, 20 mg

#### TA-01

Cat. No.: HY-100114

TA-01 is a potent CK1 and p38 MAPK inhibitor, with IC<sub>so</sub>s of 6.4 nM, 6.8 nM, 6.7 nM for CK1ε, CK1δ and p38 MAPK, respectively. TA-01 acts as a cardiogenic inhibitor.



Purity: 99.77%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

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#### TA-02

TA-02, an analog of SB 203580 (HY-10256), is a p38 MAPK inhibitor with an IC<sub>so</sub> of 20 nM. TA-02 especially inhibits TGFBR-2. TA-02 exhibits similar cardiogenic properties as SB 203580 and SB 202190 (HY-10295).

Purity: 99 57%

Clinical Data: No Development Reported

Size:



Cat. No.: HY-100115

# 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 50 mg, 100 mg

#### Tacrolimus monohydrate (FK506 monohydrate; Fujimycin monohydrate; FR900506 monohydrate) Cat. No.: HY-13756A

Tacrolimus monohydrate (FK506 monohydrate), a

macrocyclic lactone, binds to FK506 binding protein (FKBP) to form a complex and inhibits calcineurin phosphatase, which inhibits T-lymphocyte signal transduction and IL-2 transcription. Immunosuppressive properties.

99 37% **Purity:** Clinical Data: Launched

10 mM × 1 mL, 5 mg, 10 mg, 50 mg Size:



#### Talarozole

(R115866) Cat. No.: HY-14531

Talarozole (R115866) is an oral systemic all-trans retinoic acid metabolism blocking agent (RAMBA) which increases intracellular levels of endogenous all-trans retinoic acid (RA). Talarozole inhibits both CYP26A1 and CYP26B1 with ICsos of 5.4 and 0.46 nM, respectively.

99.78% Purity: Clinical Data: Phase 2

Size: 10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 50 mg, 100 mg



#### Tamoxifen

(ICI 47699; (Z)-Tamoxifen; trans-Tamoxifen) Cat. No.: HY-13757A

Tamoxifen (ICI 47699) is an orally active, selective estrogen receptor modulator (SERM) which blocks estrogen action in breast cells and can activate estrogen activity in other cells, such as bone, liver, and uterine cells.

99.92% Purity: Clinical Data: Launched

10 mM × 1 mL, 500 mg, 1 g, 5 g Size

#### Tamoxifen-d5

(ICI 47699-d5; (Z)-Tamoxifen-d5; trans-Tamoxifen-d5) Cat. No.: HY-13757AS

Tamoxifen-d5 (ICI 47699-d5) is a deuterium labeled Tamoxifen. Tamoxifen (ICI 47699) is an orally active, selective estrogen receptor modulator (SERM). Tamoxifen is a potent Hsp90 activator and enhances the Hsp90 molecular chaperone ATPase activity.

Purity: >98%

Clinical Data: No Development Reported

Size 1 mg, 5 mg

#### **Tacrolimus**

(FK506; Fujimycin; FR900506)

Tacrolimus (FK506), a macrocyclic lactone, binds to FK506 binding protein (FKBP) to form a complex. Tacrolimus inhibits calcineurin phosphatase, which inhibits T-lymphocyte signal transduction and IL-2 transcription. Immunosuppressive properties.

**Purity:** 99 93% Clinical Data: Launched

Size: 10 mg, 50 mg, 100 mg, 200 mg, 500 mg



Cat. No.: HY-13756

#### Tacrolimus-13C,d2

(FK506-13C,d2; Fujimycin-13C,d2; FR900506-13C,d2)

Tacrolimus-13C,D2 (FK506-13C,D2) is a 13C-labeled and deuterium labeled Tacrolimus, Tacrolimus (FK506), a macrocyclic lactone, binds to FK506 binding protein (FKBP) to form a complex.

**Purity:** >98%

Clinical Data: No Development Reported



Cat. No.: HY-13756S

#### **Tamibarotene**

(Am 80) Cat. No.: HY-14652

Tamibarotene is a retinoic acid receptor  $\alpha/\beta$ (RARα/β) agonist, showing high selectivity over RARy.

99 94% Purity: Clinical Data: Launched

10 mM × 1 mL, 10 mg, 50 mg Size

#### Tamoxifen Citrate (ICI 46474; (Z)-Tamoxifen Citrate;

trans-Tamoxifen Citrate) Cat. No.: HY-13757

Tamoxifen Citrate (ICI 46474) is an orally active, selective estrogen receptor modulator (SERM) which blocks estrogen action in breast cells and can activate estrogen activity in other cells, such as bone, liver, and uterine cells.

99.93% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g



#### Tanespimycin

(17-AAG; NSC 330507; CP 127374)

Tanespimycin (17-AAG) is a potent HSP90 inhibitor with an IC<sub>50</sub> of 5 nM, having a 100-fold higher binding affinity for tumour cell derived HSP90 than normal cell derived HSP90. Tanespimycin depletes cellular STK38/NDR1 and reduces STK38 kinase activity.

Purity: 99.07% Clinical Data: Phase 3

10 mM × 1 mL, 10 mg, 25 mg, 100 mg, 200 mg

Cat. No.: HY-10211

#### Tarenflurbil

((R)-Flurbiprofen; MPC7869) Cat. No.: HY-10291

Tarenflurbil ((R)-Flurbiprofen) is the R-enantiomer of the racemate NSAID Flurbiprofen, Tarenflurbil ((R)-Flurbiprofen) inhibits the binding of [ $^3$ H]9-cis-RA to RXR $\alpha$ LBD with IC $_{\!50}$  of 75  $\mu$ M. Tarenflurbil can be used for Alzheimer's disease research.

Cat. No.: HY-19934A

Purity: 99.96% Clinical Data: Phase 3

Size: 10 mM × 1 mL, 100 mg

## TAS-117

TAS-117 is a potent, selective, orally active allosteric **Akt** inhibitor (with  $\rm IC_{50}$ 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.



Cat. No.: HY-19934

**Purity:** >98%

Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg, 50 mg

# Tat-beclin 1

Cat. No.: HY-P2260

Tat-beclin 1, a peptide derived from a region of the autophagy protein (beclin 1), is a potent inducer of **autophagy** and interacts with negative regulator of autophagy, GAPR-1 (GLIPR2).

YORKRROPPROGTNYFNATFEMHDGEFG

Purity: 99.68%

Clinical Data: No Development Reported
Size: 1 mg, 5 mg, 10 mg

#### TAS-117 hydrochloride

TAS-117 hydrochloride is a potent, selective, orally active allosteric **Akt** inhibitor (with  $IC_{sq}$ s of 4.8, 1.6, and 44 nM for Akt1, 2, and 3,

respectively).

Purity: 98.96% Clinical Data: Phase 2

Size: 5 mg, 10 mg, 25 mg

#### Taurine

#### (2-Aminoethanesulfonic acid) Cat. No.: HY-B0351

Taurine, a sulphur-containing amino acid and an organic osmolyte involved in cell volume regulation, provides a substrate for the formation of bile salts, and plays a role in the modulation of intracellular free calcium concentration.



Purity: ≥98.0% Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 1 g, 5 g

#### Tat-beclin 1 TFA

Cat. No.: HY-P2260A

Tat-beclin 1 TFA, a peptide derived from a region of the autophagy protein (beclin 1), is a potent inducer of **autophagy** and interacts with negative regulator of autophagy, GAPR-1 (GLIPR2).

**Purity:** >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Taurine-13C2

## (2-Aminoethanesulfonic acid-13C2) Cat. No.: HY-B0351S1

Taurine-13C2 (2-Aminoethanesulfonic acid-13C2) is the 13C-labeled Taurine.

**Purity:** >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

### Taurine-13C2,15N

## (2-Aminoethanesulfonic acid-13C2,15N) Cat. No.: HY-B0351S2

Taurine-13C2,15N (2-Aminoethanesulfonic acid-13C2,15N) is the 13C- and 15N- labeled

Taurine.

Cat. No.: HY-N0136

**Purity:** >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Taurine-d4

#### (2-Aminoethanesulfonic acid-d4) Cat. No.: HY-B0351S

Taurine-d4 (2-Aminoethanesulfonic acid-d4) is the deuterium labeled Taurine.

**Purity:** > 98%

Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg

#### Taxifolin

#### ((+)-Dihydroquercetin; (+)-Taxifolin)

Taxifolin ((+)-Dihydroquercetin) exhibits important anti-tyrosinase activity. Taxifolin exhibits significant inhibitory activity against collagenase with an  $\rm IC_{50}$  value of 193.3  $\mu M$ . Taxifolin is an important natural compound with antifibrotic activity.

with HO OH OH

**Purity:** 99.97%

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

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#### Taxifolin-d3

((+)-Dihydroquercetin-d3; (+)-Taxifolin-d3)

Taxifolin-d3 is deuterium labeled Taxifolin. Taxifolin ((+)-Dihydroguercetin) exhibits important anti-tyrosinase activity. Taxifolin exhibits significant inhibitory activity against collagenase with an IC50 value of 193.3  $\mu M$ .

Cat. No.: HY-N0136S

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### **TBHQ**

Purity:

Size:

Tazarotene (AGN 190168)

#### (tert-Butylhydroquinone)

Clinical Data: Launched

TBHQ (tert-Butylhydroguinone) is a widely used Nrf2 activator, protects against Doxorubicin (DOX)-induced cardiotoxicity through activation of

10 mM × 1 mL, 10 mg, 50 mg, 100 mg

Tazarotene (AGN 190168) is a selective retinoic

plaque psoriasis and acne vulgaris.

99 93%

acid receptor (RAR) agonist for the treatment of

Nrf2



Cat. No.: HY-12248

ale music

Cat. No.: HY-100489

Cat. No.: HY-15388

**Purity:** 99 76%

Telaglenastat (CB-839)

C) compared to GLS2.

Clinical Data: Phase 2

Clinical Data: No Development Reported

10 mM × 1 mL, 500 mg, 1 g

Telaglenastat (CB-839) is a first-in-class,

glutaminase 1 (GLS1) inhibitor. Telaglenastat

(kidney-type glutaminase) and GAC (glutaminase

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

selectively inhibits GLS1 splice variants KGA

selective, reversible and orally active

99.82%

#### Tazarotene-d8

Cat. No.: HY-15388S

Tazarotene-d8 is the deuterium labeled Tazarotene. Tazarotene (AGN 190168) is a selective retinoic acid receptor (RAR) agonist for the treatment of plaque psoriasis and acne vulgaris.

Purity: >98%

Clinical Data: No Development Reported

1 mg, 10 mg

## Tea polyphenol

Cat. No.: HY-N1925

Tea polyphenol is the floorboard of phenolic compounds in tea. Tea polyphenol exhibits biological activity including antioxidant and anti-cancer activities, inhibition of cell proliferation, induction of apoptosis, cell cycle arrest and modulation of carcinogen metabolism.

Tea polyphenol

≥99.0% Purity: Clinical Data: Phase 3 Size: 100 ma

#### Telaglenastat hydrochloride

(CB-839 hydrochloride) Cat. No.: HY-12248A

Telaglenastat (CB-839) hydrochloride is a first-in-class, selective, reversible and orally active glutaminase 1 (GLS1) inhibitor.

against a

>98% Purity:

Clinical Data: No Development Reported

Size: 1 ma, 5 ma

#### Telmisartan

(BIBR 277)

**Purity:** 

Size

Telmisartan is a potent, long lasting antagonist of angiotensin II type 1 receptor (AT1), selectively inhibiting the binding of 125I-AngII to AT1 receptors with IC<sub>so</sub> of 9.2 nM.

Cat. No.: HY-13955

99.96% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 50 mg, 100 mg, 500 mg, 1 g

#### Telmisartan-13C,d3

(BIBR 277-13C,d3) Cat. No.: HY-13955S2

Telmisartan-13C,d3 is the 13C- and deuterium labeled. Telmisartan is a potent, long lasting antagonist of angiotensin II type 1 receptor (AT1), selectively inhibiting the binding of 125I-AngII to AT1 receptors with IC50 of 9.2 nM.



Purity: >98%

Clinical Data: No Development Reported

Size 1 mg, 5 mg

#### Telmisartan-d3

Telmisartan-d3 is the deuterium labeled Telmisartan. Telmisartan is a potent, long lasting antagonist of angiotensin II type 1 receptor

(AT1), selectively inhibiting the binding of <sup>125</sup>I-AngII to AT1 receptors with IC<sub>50</sub> of 9.2 nM.

Purity: >98%

Clinical Data: No Development Reported

1 mg, 10 mg



**123** 

Cat. No.: HY-13955S

#### Telmisartan-d4

Cat. No.: HY-13955S1

Telmisartan-d4 is the deuterium labeled Telmisartan, Telmisartan is a potent, long lasting antagonist of angiotensin II type 1 receptor (AT1), selectively inhibiting the binding of <sup>125</sup>I-AngII to AT1 receptors with IC<sub>50</sub> of 9.2 nM.



>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Temozolomide-d3

Temozolomide-d3 (NSC 362856-d3) is the deuterium labeled Temozolomide. Temozolomide (NSC 362856) is an oral active DNA alkylating agent that crosses the blood-brain barrier. Temozolomide is also a proautophagic and proapoptotic agent.



Cat. No.: HY-17364S

**Purity:** >98%

Clinical Data:

1 mg, 5 mg

#### Tempol

Purity:

Size:

(4-Hydroxy-TEMPO)

Clinical Data: Launched

Temozolomide

proapoptotic agent.

(NSC 362856; CCRG 81045; TMZ)

99 98%

Temozolomide (NSC 362856) is an oral active DNA

barrier. Temozolomide is also a proautophagic and

alkylating agent that crosses the blood-brain

Tempol is a general superoxide dismutase (SOD)-mimetic drug that efficiently neutralizes reactive oxygen species (ROS).



Cat. No.: HY-100561

Cat. No.: HY-17364

**Purity:** 99 98% Clinical Data: Phase 2

10 mM × 1 mL, 200 mg, 1 g

#### **Temsirolimus**

(CCI-779) Cat. No.: HY-50910

Temsirolimus is an inhibitor of mTOR with an IC<sub>50</sub> of 1.76 μM. Temsirolimus activates autophagy and prevents deterioration of cardiac function in animal model.



99 56% Purity: Clinical Data: Launched

Size:  $10 \text{ mM} \times 1 \text{ mL}$ , 5 mg, 10 mg, 25 mg, 100 mg

## Temsirolimus-d3

(CCI-779-d3) Cat. No.: HY-50910S

10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg, 500 mg, 1 g

Temsirolimus-d3 (CCI-779-d3) is the deuterium labeled Temsirolimus. Temsirolimus is an inhibitor of mTOR with an  $IC_{so}$  of 1.76  $\mu$ M. Temsirolimus activates autophagy and prevents deterioration of cardiac function in animal model.



>98% Purity:

Clinical Data: No Development Reported

Size 1 mg, 5 mg

#### Temsirolimus-d3-1

(CCI-779-d3-1) Cat. No.: HY-50910S2

Temsirolimus-d3-1 (CCI-779-d3-1) is the deuterium labeled Temsirolimus. Temsirolimus is an inhibitor of mTOR with an  $IC_{so}$  of 1.76  $\mu$ M. Temsirolimus activates autophagy and prevents deterioration of cardiac function in animal model.



Purity: >98%

Clinical Data: No Development Reported

Size 1 mg, 5 mg

#### Temsirolimus-d7

(CCI-779-d7) Cat. No.: HY-50910S1

Temsirolimus-d7 (CCI-779-d7) is the deuterium labeled Temsirolimus. Temsirolimus is an inhibitor of mTOR with an  $IC_{so}$  of 1.76  $\mu$ M. Temsirolimus activates autophagy and prevents deterioration of cardiac function in animal model.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Tenovin-6

Cat. No.: HY-15510

Tenovin-6, an analog of Tenovin-1 (HY-13423), is an activator of p53 transcriptional activity. Tenovin-6 inhibits the protein deacetylase activities of purified human SIRT1, SIRT2, and SIRT3 with IC  $_{50}$ s of 21  $\mu$ M, 10  $\mu$ M, and 67  $\mu$ M, respectively.



**Purity:** 98.67%

Clinical Data: No Development Reported

10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 50 mg, 100 mg Size:

#### Tenovin-1

Cat. No.: HY-13423

Tenovin-1, a p53 activator, protects p53 from MDM2-mediated degradation. Tenovin-1 acts through inhibition of the protein-deacetylating activities of SirT1 and SirT2. Tenovin-1 is also a dihydroorotate dehydrogenase (DHODH) inhibitor.



Purity: 99.88%

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

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#### Tenovin-6 Hydrochloride

Cat. No.: HY-15510B

Tenovin-6 Hydrochloride, an analog of Tenovin-1 (HY-13423), is an activator of p53 transcriptional activity.

>98.0% Purity:

(HZIV 81-2 D6)

Tetrahydrocurcumin D6

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 50 mg, 100 mg

Tetrahydrocurcumin D6 (HZIV 81-2 D6) is a deuterium labeled Tetrahydrocurcumin. Tetrahydrocurcumin is a Curcuminoid which displays inhibitory activity for CYP2C9 and CYP3A4.

Cat. No.: HY-N0893S

Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg

# TFEB activator 1

TFEB activator 1 is an orally effective, mTOR-independent activator of TFEB. TFEB activator 1 significantly promotes the nuclear translocation of Flag-TFEB with an EC<sub>so</sub> of 2167 nM.

Cat. No.: HY-14658S

Cat. No.: HY-135825

Purity: 99.69%

Clinical Data: No Development Reported

Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

#### Thalidomide D4

Thalidomide D4 is a deuterium labeled Thalidomide. Thalidomide inhibits cereblon (CRBN), a part of the cullin-4 E3 ubiquitin ligase complex CUL4-RBX1-DDB1, with a K<sub>d</sub> of ~250 nM, and has immunomodulatory, anti-inflammatory and anti-angiogenic cancer properties.

Clinical Data: No Development Reported

Size:

# 5 mg, 10 mg

#### Theophylline-d6

#### (1,3-Dimethylxanthine-d6; Theo-24-d6) Cat. No.: HY-B0809S

Theophylline-d6 (1,3-Dimethylxanthine-d6) is the deuterium labeled Theophylline. Theophylline is a nonselective phosphodiesterase (PDE) inhibitor, adenosine receptor blocker, and histone deacetylase (HDAC) activator.

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Tetrahydrocurcumin

(HZIV 81-2) Cat. No.: HY-N0893

Tetrahydrocurcumin is a Curcuminoid found in turmeric (Curcuma longa) that is produced by the reduction of Curcumin. Tetrahydrocurcumin inhibit CYP2C9 and CYP3A4.



≥95.0% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

#### Tezacaftor

(VX-661) Cat. No.: HY-15448

Tezacaftor (VX-661) is a second F508del CFTR corrector and help CFTR protein reach the cell surface.



**Purity:** 99 94% Clinical Data: Launched

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

#### TG101209

TG101209 is a selective JAK2 inhibitor with IC<sub>50</sub> of 6 nM, less potent to Flt3 and RET with IC<sub>so</sub> of 25 nM and 17 nM, appr 30-fold selective for JAK2 than JAK3, and sensitive to JAK2V617F and MPLW515L/K mutations.



Cat. No.: HY-10410

99.72% Purity:

Clinical Data: No Development Reported

Size 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

#### Theophylline

## (1,3-Dimethylxanthine; Theo-24)

Theophylline is a nonselective phosphodiesterase (PDE) inhibitor, adenosine receptor blocker, and histone deacetylase (HDAC) activator.



Cat. No.: HY-B0809

99.94% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 500 mg, 5 g

#### Thiamet G

Thiamet G is a potent and selective inhibitor of O-GlcNAcase (OGA), which acts to remove O-GlcNAc from modified proteins, with K, of 20 nM for human OGA.



Cat. No.: HY-12588

99.98%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg

#### Thioridazine

Thioridazine, an antagonist of the dopamine 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: Launched
Size: 1 mg, 5 mg



Cat. No.: HY-B0965A

#### Thioridazine hydrochloride

Thioridazine hydrochloride, an orally active antagonist of the **dopamine receptor D2** family proteins, exhibits potent anti-psychotic and anti-anxiety activities.

Purity: 99.93% Clinical Data: Launched

Size: 10 mM × 1 mL, 100 mg, 500 mg



Cat. No.: HY-B0965

H-CI

#### Thioridazine-d3 2-Sulfone

Thioridazine-d3 2-Sulfone is the deuterium labeled Thioridazine hydrochloride. Thioridazine hydrochloride, an orally active antagonist of the **dopamine receptor D2** family proteins, exhibits potent anti-psychotic and anti-anxiety activities.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 10 mg

D N O

Cat. No.: HY-B0965S

#### Thioridazine-d3 hydrochloride

Cat. No.: HY-B0965AS

Thioridazine-d3 hydrochloride is the deuterium labeled Thioridazine. Thioridazine, an antagonist of the dopamine receptor D2 family proteins, exhibits potent anti-psychotic and anti-anxiety activities.

dividies.

**Purity:** 

>98%

Clinical Data: No Development Reported

Size: 1 mg, 10 mg



## THZ-P1-2

Cat. No.: HY-136351

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THZ-P1-2 is a first-in-class and selective PI5P4K inhibitor, with an IC $_{50}$  of 190 nM for PI5P4K $\alpha$ . THZ-P1-2 covalently targets cysteines on a disordered loop in PI5P4K $\alpha/\beta/\gamma$ . THZ-P1-2 causes autophagy disruption and upregulates TFEB signaling.

Purity: 99.06%

Clinical Data: No Development Reported
Size: 5 mg, 10 mg, 50 mg, 100 mg

## Tigecycline

(GAR-936) Cat. No.: HY-B0117

Tigecycline (GAR-936) is a broad-spectrum glycylcycline antibiotic. The mean inhibitory concentration (MIC) of Tigecycline for E. coli (MG1655 strain) is approximately 125 ng/mL.



Purity: 99.74% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg, 500 mg

#### Tigecycline hydrate

(GAR-936 hydrate) Cat. No.: HY-B0117D

Tigecycline hydrate (GAR-936 hydrate) is a broad spectrum glycylcycline antibiotic.

Purity: >98%
Clinical Data: Phase 4
Size: 1 mg, 5 mg

#### Tigecycline hydrochloride

(GAR-936 hydrochloride)

Tigecycline hydrochloride (GAR-936 hydrochloride) is a broad-spectrum glycylcycline antibiotic. The mean inhibitory concentration (MIC) of Tigecycline for E. coli (MG1655 strain) is approximately 125 ng/mL.

Cat. No.: HY-B0117A

Purity: >98%
Clinical Data: Launched
Size: 1 mg, 5 mg

# Tigecycline mesylate

(GAR-936 mesylate) Cat. No.: HY-B0117B

Tigecycline mesylate (GAR-936 mesylate) is a broad-spectrum glycylcycline antibiotic. The mean inhibitory concentration (MIC) of Tigecycline for E. coli (MG1655 strain) is approximately 125 ng/mL.



Purity: >98%
Clinical Data: Launched
Size: 1 mg, 5 mg

#### Tigecycline tetramesylate

(GAR-936 tetramesylate)

Tigecycline tetramesylate (GAR-936 tetramesylate) is a broad-spectrum glycylcycline antibiotic. The mean inhibitory concentration (MIC) of Tigecycline for E. coli (MG1655 strain) is approximately 125

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Cat. No.: HY-B0117C

Purity: 95.36% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg

#### Tigecycline-d9

(GAR-936-d9) Cat. No.: HY-B0117S

Tigecycline-d9 is deuterium labeled Tigecycline. Tigecycline (GAR-936) is a broad-spectrum glycylcycline antibiotic. The mean inhibitory concentration (MIC) of Tigecycline for E. coli (MG1655 strain) is approximately 125 ng/mL.



>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### **Tolbutamide**

Purity:

Size:

Tolbutamide is a first generation potassium channel blocker, sulfonylurea oral hypoglycemic drug. Target: Potassium Channel Tolbutamide is an oral antihyperglycemic agent used for the treatment of non-insulin-dependent diabetes mellitus (NIDDM).

**Purity:** 99 96% Clinical Data: Launched

10 mM × 1 mL, 500 mg, 1 g, 5 g

Tizoxanide D4

Tizoxanide D4 (TIZ D4) is the deuterium labeled Tizoxanide Tizoxanide is the active metabolite of Nitazoxanide, which is a thiazolide anti-infective compound against anaerobic bacteria, protozoa, and a range of viruses. Tizoxanide has anti-HIV-1 activities.

Purity: >98%

Clinical Data: No Development Reported

1 mg, 5 mg



Cat. No.: HY-12687S

#### Tolbutamide-d9

Cat. No.: HY-B0401S

Tolbutamide-d9 is the deuterium labeled Tolbutamide. Tolbutamide is a first generation potassium channel blocker, sulfonylurea oral hypoglycemic drug.

Purity: >98%

Tolvaptan-D7

Clinical Data: No Development Reported

Size: 2.5 mg, 25 mg

## Tolvaptan

(OPC-41061)

Tolvaptan is a selective, competitive arginine vasopressin receptor 2 antagonist with an IC50 of 1.28µM for the inhibition of AVP-induced platelet aggregation.

Purity: 99.96% Clinical Data: Launched

Size 10 mM × 1 mL, 10 mg, 50 mg

**Tomatidine** 

# Cat. No.: HY-17000S

Tolvaptan-D7 (OPC-41061-D7) is the deuterium labeled Tolvaptan. Tolvaptan is a selective, competitive arginine vasopressin receptor 2 antagonist with an  $IC_{so}$  of 1.28 $\mu$ M for the inhibition of AVP-induced platelet aggregation.



>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

activates autophagy either in mammal cells or C elegans.

≥95.0% Purity:

Clinical Data: No Development Reported

Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

Tomatidine acts as an anti-inflammatory agent by

blocking NF-κB and JNK signaling. Tomatidine

#### Tomatidine hydrochloride

Cat. No.: HY-N2149A

Tomatidine hydrochloride acts as an anti-inflammatory agent by blocking NF-κB and JNK signaling. Tomatidine hydrochloride activates autophagy either in mammal cells or C elegans.



Purity: ≥98.0%

Clinical Data: No Development Reported 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

#### Topotecan

(SKF 104864A; NSC 609669)

Topotecan (SKF 104864A; NSC 609669) is a Topoisomerase I inhibitor. The IC<sub>50</sub> values of Topotecan at 24 h are 2.73±0.25 µM of U251 cells,  $2.95\pm0.23~\mu\text{M}$  of U87 cells,  $5.46\pm0.41~\mu\text{M}$  of

GSCs-U251 and  $5.95\pm0.24~\mu M$  of GSCs-U87. >98%

Purity: Clinical Data: Launched

10 mg, 50 mg, 100 mg



Tizoxanide is the active metabolite of

98 10%

Clinical Data: No Development Reported

Nitazoxanide, which is a thiazolide anti-infective

a range of viruses. Tizoxanide has anti-HIV-1

compound against anaerobic bacteria, protozoa, and

10 mM × 1 mL, 10 mg, 50 mg, 100 mg

(TIZ) Cat. No.: HY-12687



Cat. No.: HY-B0401

Cat. No.: HY-17000

Cat. No.: HY-N2149

Cat. No.: HY-13768

#### Topotecan Hydrochloride

(SKF 104864A Hydrochloride; NSC 609669 Hydrochloride) Cat. No.: HY-13768A

Topotecan Hydrochloride (SKF 104864A Hydrochloride) is a **Topoisomerase I** inhibitor with potent antineoplastic activities.

Purity: 99.74% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg

#### Topotecan-d5

Topotecan-d5 is the deuterium labeled Topotecan. Topotecan (SKF 104864A; NSC 609669) is a **Topoisomerase I** inhibitor. The  $IC_{so}$  values of Topotecan at 24 h are 2.73±0.25  $\mu$ M of U251 cells, 2.95±0.23  $\mu$ M of U87 cells, 5.46±0.41  $\mu$ M of GSCs-U251 and 5.95±0.24  $\mu$ M of GSCs-U87.

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Cat. No.: HY-13768S

**Purity:** >98%

Clinical Data: No Development Reported Size: 1 mg, 10 mg, 50 mg

#### Topotecan-d6

Cat. No.: HY-13768S1

Topotecan-d6 is the deuterium labeled Topotecan. Topotecan (SKF 104864A; NSC 609669) is a **Topoisomerase** I inhibitor. The  $IC_{50}$  values of Topotecan at 24 h are 2.73±0.25  $\mu M$  of U251 cells, 2.95±0.23  $\mu M$  of U87 cells, 5.46±0.41  $\mu M$  of GSCs-U251 and 5.95  $\mu M$  of GSCs-U87.



**Purity:** >98%

Clinical Data: No Development Reported Size: 1 mg, 5 mg, 10 mg

#### Torin 1

Torin 1 is a potent inhibitor of mTOR with an  $IC_{50}$  of 3 nM. Torin 1 inhibits both mTORC1/2 complexes with  $IC_{50}$  values between 2 and 10 nM. Torin 1 is an effective inducer of autophagy.

Cat. No.: HY-13003

**Purity:** 98.95%

Clinical Data: No Development Reported Size: 5 mg, 10 mg, 50 mg, 100 mg

#### Torin 2

Cat. No.: HY-13002

Torin 2 is an mTOR inhibitor with EC $_{50}$  of 0.25 nM for inhibiting cellular mTOR activity, and exhibits 800-fold selectivity over PI3K (EC $_{50}$ : 200 nM). Torin 2 also inhibits DNA-PK with an IC $_{50}$  of 0.5 nM in the cell free assay. Torin 2 can suppress both mTORC1 and mTORC2.



**Purity:** 99.98%

Clinical Data: No Development Reported

Size:  $10 \text{ mM} \times 1 \text{ mL}$ , 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

#### Torkinib

(PP 242)

Torkinib (PP 242) is a selective and ATP-competitive mTOR inhibitor with an  $IC_{50}$  of 8 nM. PP242 inhibits both mTORC1 and mTORC2 with  $IC_{50}$  of 30 nM and 58 nM, respectively.



Cat. No.: HY-10474

**Purity:** 98.76%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

#### Tozasertib

(VX 680; MK-0457) Cat. No.: HY-10161

Tozasertib (VX 680; MK-0457) is an inhibitor of Aurora A/B/C kinases with  $\rm K_i s$  of 0.6, 18, 4.6 nM, respectively.



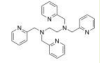
Purity: 99.94% Clinical Data: Phase 2

Size:  $10 \text{ mM} \times 1 \text{ mL}$ , 50 mg, 100 mg, 250 mg

#### **TPEN**

(TPEDA) Cat. No.: HY-100202

TPEN (TPEDA) is a specific cell-permeable heavy metal chelator. TPEN has a higher affinity for Zn²+, but a lower affinity for Mg²+ and Ca²+. TPEN induces DNA damage and increases intracellular ROS production. TPEN also inhibits cell proliferation and induces apoptosis.



**Purity:** 99.21%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 50 mg, 100 mg, 200 mg

#### Trametinib

(GSK1120212; JTP-74057) Cat. No.: HY-10999

Trametinib (GSK1120212; JTP-74057) is an orally active **MEK** inhibitor that inhibits MEK1 and MEK2 with  $IC_{50}$ s of about 2 nM. Trametinib activates **autophagy** and induces **apoptosis**.



Purity: 99.92% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg

#### Trametinib-13C,d3

(GSK1120212-13C,d3; JTP-74057-13C,d3)

Trametinib-13C,d3 is the 13C- and deuterium labeled. Trametinib (GSK1120212; JTP-74057) is an orally active MEK inhibitor that inhibits MEK1 and MEK2 with IC50s of about 2 nM. Trametinib activates autophagy and induces apoptosis.



Cat. No.: HY-10999S2

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

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

#### Trametinib-13C6

Cat. No.: HY-10999S1

Trametinib-13C6 is the 13C-labeled Trametinib. Trametinib (GSK1120212: JTP-74057) is an orally active MEK inhibitor that inhibits MEK1 and MEK2 with IC<sub>50</sub>s of about 2 nM. Trametinib activates autophagy and induces apoptosis.

Purity: >98%

Clinical Data: No Development Reported

Size:

# 1 mg, 5 mg

## Trametinib-d4

Trametinib-d4 is the deuterium labeled Trametinib. Trametinib (GSK1120212: JTP-74057) is an orally active MEK inhibitor that inhibits MEK1 and MEK2 with IC<sub>50</sub>s of about 2 nM. Trametinib activates autophagy and induces apoptosis.

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg



Cat. No.: HY-10999S

#### Triclosan

Cat. No.: HY-B1119

Triclosan is an antibacterial and antifungal agent found in consumer products, including soaps, detergents, toys, and surgical cleaning treatments.

≥97.0% Purity: Clinical Data: Launched

10 mM × 1 mL, 100 mg

#### Triclosan-d3

Triclosan D3 is the deuterium labeled Triclosan. Triclosan is an antibacterial and antifungal agent found in consumer products, including soaps, detergents, toys, and surgical cleaning

treatments.

**Purity:** >98%

Clinical Data: No Development Reported

1 mg, 5 mg



Cat. No.: HY-B1119S

Trifarotene

(CD5789) Cat. No.: HY-100256

Trifarotene (CD5789) is a potent and selective RARy agonist. Trifarotene (CD5789) shows 65-fold and 16-fold selectivitiy for the RARy (EC $_{sn}$ =7.7 nM) over RAR $\alpha$  (EC $_{s0}$ =500 nM) and RARβ ( $EC_{so}$ =125 nM), respectively.

Purity: 99.50% Clinical Data: Launched

Size:  $10 \text{ mM} \times 1 \text{ mL}$ , 5 mg, 10 mg, 50 mg, 100 mg

#### Triflupromazine hydrochloride

Triflupromazine hydrochloride is an antipsychotic medication, which are Dopamine D1/D2 receptor

antagonists.

H-CI

Cat. No.: HY-B0909

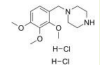
99.80% Purity: Clinical Data: Launched

Size 10 mM × 1 mL, 100 mg

# Trimetazidine dihydrochloride

Cat. No.: HY-B0968

Trimetazidine dihydrochloride is a selective long chain 3-ketoyl coenzyme A thiolase inhibitor with an  $IC_{50}$  of 75 nM, which can inhibit  $\beta$ -oxidation of free fatty acid (FFA).



Cat. No.: HY-13067

99.62% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg

# Trimetazidine

Trimetazidine is a selective long chain 3-ketoyl

coenzyme A thiolase inhibitor with an IC<sub>50</sub> of 75 nM, which can inhibit  $\beta\text{-}oxidation$  of free fatty acid (FFA).

Cat. No.: HY-B0968A

99.55% Purity: Clinical Data: Launched Size: 10 mg, 50 mg

#### Trimetazidine-d8 dihydrochloride

Cat. No.: HY-B0968S

Trimetazidine-d8 dihydrochloride is the deuterium labeled Trimetazidine dihydrochloride. Trimetazidine dihydrochloride is a selective long chain 3-ketoyl coenzyme A thiolase inhibitor with an  $IC_{so}$  of 75 nM, which can inhibit  $\beta$ -oxidation of free fatty acid (FFA).

HCI HCI

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 10 mg

#### Tripterin (Celastrol)

Tripterin (Celastrol) is a proteasome inhibitor which potently and preferentially inhibits the chymotrypsin-like activity of a purified 20S proteasome with  $IC_{50}$  of 2.5  $\mu$ M.

Purity: 99.90%

Clinical Data: No Development Reported

10 mM × 1 mL, 10 mg, 50 mg, 100 mg

#### **Triptonide**

(NSC 165677; PG 492) Cat. No.: HY-32736

Triptonide (NSC 165677) is a natural product identified in Tripterygium wilfordii Hook F.. Triptonide is a **Wnt** signaling inhibitor with an  $IC_{50}$  of appropriately 0.3nM.



Purity: 99.73% Clinical Data: Phase 2

Size: 10 mM × 1 mL, 10 mg

## Troglitazone

(CS-045) Cat. No.: HY-50935

Troglitazone is a PPAR $\gamma$  agonist, with EC $_{50}$ S of 550 nM and 780 nM for human and murinePPAR $\gamma$  receptor, respectively.



Purity: 98.60% Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg

## Troglitazone-d4

(CS-045-d4) Cat. No.: HY-50935S

Troglitazone-d4 is deuterium labeled Troglitazone.
Troglitazone is a PPARy agonist, with EC50s of 550 nM and 780 nM for human and murinePPARy receptor, respectively.

**Purity:** >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Tropifexor

(LJN452) Cat. No.: HY-107418

Tropifexor (LJN452) is a highly potent agonist of FXR with an EC<sub>so</sub> of 0.2 nM.



Purity: 99.35% Clinical Data: Phase 2

Size: 10 mM × 1 mL, 1 mg, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

#### **TTNPB**

(Ro 13-7410; Arotinoid acid; AGN191183) Cat. No.: HY-15682

TTNPB is a highly potent RAR agonist. Competitive binding assays using human RARs yield  $IC_{s0}s$  of  $\alpha{=}5.1$  nM,  $\beta{=}$  4.5 nM, and  $\gamma{=}9.3$  nM, respectively.

**Purity:** 98.81%

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 5 mq, 10 mq, 50 mq

#### Tubastatin A

Cat. No.: HY-13271A

Tubastatin A is a potent and selective HDAC6 inhibitor with an  $\rm IC_{50}$  of 15 nM in a cell-free assay, and is selective (1000-fold more) against all other isozymes except HDAC8 (57-fold more).



**Purity:** 98.12%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg

#### Tubastatin A Hydrochloride

(Tubastatin A HCl; TSA HCl)

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).



Cat. No.: HY-13271

Purity: 98.21%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

#### Turofexorate isopropyl

(FXR-450; XL335; WAY-362450)

Turofexorate isopropyl (FXR-450) is a potent, selective, and orally bioavailable FXR agonist with EC $_{\rm sn}$  of 4 nM.



Cat. No.: HY-50911

Purity: 99.63% Clinical Data: Phase 1

Size:  $10 \text{ mM} \times 1 \text{ mL}$ , 5 mg, 10 mg, 50 mg, 100 mg

#### TWS119

Cat. No.: HY-10590

TWS119 is a specific inhibitor of GSK-3 $\beta$ , with an IC $_{50}$  of 30 nM, and activates the wnt/ $\beta$ -catenin pathway.

**Purity:** ≥98.0%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 10 mg, 50 mg, 100 mg

#### Typhaneoside

Typhaneoside, extracted from Typha angustifolia L., Typhaneoside can inhibit the excessive **autophagy** of hypoxia/reoxygenation cells and increase the phosphorylation of Akt and mTOR.



Cat. No.: HY-N0712

**Purity:** 99.74%

Clinical Data: No Development Reported
Size: 5 mg, 10 mg, 20 mg

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#### U0126

U0126 is a potent, non-ATP competitive and selective MEK1 and MEK2 inhibitor, with

IC<sub>so</sub>s of 72 nM and 58 nM, respectively. U0126 is an autophagy and mitophagy inhibitor.



Cat. No.: HY-12031A

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### U0126-EtOH

U0126 (U0126-EtOH) is a potent, non-ATP competitive and selective MEK1 and MEK2 inhibitor, with IC<sub>50</sub>s of 72 nM and 58 nM, respectively. U0126 is an autophagy and mitophagy inhibitor.



Cat. No.: HY-12031

99.41% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg, 500 mg Size:

#### UBCS039

Cat. No.: HY-115453

UBCS039 is the first synthetic, specific Sirtuin 6 (SIRT6) activator, inducing autophagy in human tumor cells, with an  $EC_{50}$  of 38  $\mu$ M.



Purity: 98 80%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

## Ulipristal acetate

(CDB-2914)

Ulipristal acetate (CDB-2914) is an orally active, selective progesterone receptor modulator (SPRM). Ulipristal acetate stimulates the autophagic response selectively in leiomyoma cells.



Cat. No.: HY-16508

**Purity:** 99 93% Clinical Data: Launched

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

#### Ulipristal acetate-d6

(CDB-2914-d6) Cat. No.: HY-16508S

Ulipristal acetate-d6 is deuterium labeled Ulipristal acetate. Ulipristal acetate (CDB-2914) is an orally active, selective progesterone receptor modulator (SPRM). Ulipristal acetate stimulates the autophagic response selectively in leiomyoma cells.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

# ULK1-IN-2

ULK1-IN-2 (compound 3s) is a potent ULK1 inhibitor. ULK1-IN-2 shows highest cytotoxic effect against cancer cell lines, with IC of 1.94 μM in A549. ULK1-IN-2 can induce apoptosis and simultaneously block autophagy, and can be used to study NSCLC (Non-small cell lung cancer).



Cat. No.: HY-143466

**Purity:** >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

# UM-164

(DAS-DFGO-II) Cat. No.: HY-112182

UM-164 (DAS-DFGO-II) is a highly potent inhibitor of c-Src with a K<sub>d</sub> of 2.7 nM. UM-164 also potently inhibits  $p38\alpha$  and  $p38\beta$ .



Purity: 98.91%

Clinical Data: No Development Reported

Size: 5 mg, 10 mg

#### UNBS5162

UNBS5162 is a pan-antagonist of CXCL chemokine expression, with anti-tumor activity.



Cat. No.: HY-16509

Purity: 99.92%

Clinical Data: No Development Reported

10 mM × 1 mL, 10 mg, 50 mg, 100 mg Size



#### UNC0638

Cat. No.: HY-15273

UNC0638 selectively inhibits G9a and GLP histone methyltransferase activity with IC<sub>50</sub>s of less than 15 nM and 19 nM, respectively. UNC0638 has anti-FMDV (foot-and-mouth disease virus) and anti-VSV (vesicular stomatitis virus) activities.



Purity: 99.73%

Clinical Data: No Development Reported

Size 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

#### UNC1999

UNC1999 is a SAM-competitive, potent and selective inhibitor of EZH2/1 with IC<sub>so</sub>s of <10 nM and 45 nM, repectively.



Cat. No.: HY-15646

99.85% **Purity:** 

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

#### UNC2400

UNC2400 is a close analog of UNC1999 with >1,000-fold lower potency than UNC1999 as a negative control for cell-based studies.



>98% Purity:

Clinical Data: No Development Reported

Size: 5 mg

Cat. No.: HY-12845

#### **URMC-099**

Cat. No.: HY-12599

URMC-099 is an orally bioavailable and potent mixed lineage kinase type 3 (MLK3) (IC<sub>50</sub>=14 nM) inhibitor with with excellent blood-brain barrier penetration properties.



Purity: 99 90%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Urolithin A, a gut-microbial metabolite of ellagic acid, exerts anti-inflammatory, antiproliferative, and antioxidant properties. Urolithin A induces autophagy and apoptosis, suppresses cell cycle progression, and inhibits DNA synthesis.

URB-597 (KDS-4103) is an orally bioavailable and

selective FAAH inhibitor, URB-597 inhibits FAAH

activity with an IC<sub>50</sub>s of approximately 5 nM in rat brain membranes, 0.5 nM in intact rat neurons,

3 nM in human liver microsomes. Antidepressant-like effects. Analgesic activity.

99.01%

Clinical Data: No Development Reported

Cat. No.: HY-100599

Cat. No.: HY-10864

**Purity:** 98.05%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

10 mM × 1 mL, 10 mg, 25 mg, 50 mg, 100 mg

#### Ursolic acid

(Prunol; Urson; Malol) Cat. No.: HY-N0140

Ursolic acid (Prunol) is a natural pentacyclic triterpenoid carboxylic acid, exerts anti-tumor effects and is an effective compound for cancer prevention and therapy.



99.66% Purity: Clinical Data: Phase 2

Size: 10 mM × 1 mL, 50 mg, 100 mg, 200 mg

#### UVI 3003

**URB-597** 

(KDS-4103)

Purity:

Size:

Urolithin A

UVI 3003 is a highly selective antagonist of retinoid X receptor (RXR), and inhibits xenopus and human  $RXR\alpha$  in Cos7 cells, with  $IC_{so}s$  of 0.22 and 0.24 µM, respectively.

Cat. No.: HY-107500

99.77% Purity:

Clinical Data: No Development Reported

Size 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg

#### Vacuolin-1

Cat. No.: HY-118630

Vacuolin-1 is a potent and cell-permeable lysosomal exocytosis inhibitor. Vacuolin-1 blocks the Ca<sup>2+</sup>-dependent exocytosis of lysosomes and prevents the release of lysosomal content without affecting the process of resealing.



Purity: 98.82%

Clinical Data: No Development Reported

Size: 5 mg, 10 mg

# Valinomycin

(NSC 122023)

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.



Cat. No.: HY-N6693

99.05% Purity:

Clinical Data: No Development Reported 10 mM × 1 mL, 5 mg, 10 mg Size

#### Valproic acid

(VPA; 2-Propylpentanoic Acid) Cat. No.: HY-10585

Valproic acid (VPA; 2-Propylpentanoic Acid) is an  $\mbox{HDAC}$  inhibitor, with  $\mbox{IC}_{\mbox{\scriptsize 50}}$  in the range of 0.5 and 2 mM, also inhibits **HDAC1** ( $IC_{50}$ , 400  $\mu$ M), and induces proteasomal degradation of HDAC2.



Purity: ≥98.0% Clinical Data: Launched

Size: 500 mg, 1 g, 5 g, 25 g

#### Valproic acid sodium

(Sodium Valproate sodium)

Valproic acid sodium salt (Sodium Valproate) is an  $\mbox{HDAC}$  inhibitor, with  $\mbox{IC}_{\mbox{\scriptsize 50}}$  in the range of 0.5 and 2 mM, also inhibits HDAC1 (IC<sub>so</sub>, 400  $\mu$ M), and induces proteasomal degradation of HDAC2.



Cat. No.: HY-10585A

≥98.0% Purity: Clinical Data: Launched

500 mg, 1 g, 5 g, 25 g

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#### Valproic acid-d14 sodium

(Sodium Valproate-d14 sodium)

Valproic acid-d14 (sodium) is deuterium labeled Valproic acid (sodium). Valproic acid sodium salt (Sodium Valproate) is an HDAC inhibitor, with IC50 in the range of 0.5 and 2 mM, also inhibits HDAC1 (IC50, 400  $\mu$ M), and induces proteasomal degradation of HDAC2.

Cat. No.: HY-10585AS1

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Valproic acid-d4

(VPA-d4; 2-Propylpentanoic Acid-d4)

Valproic acid-d4 (VPA-d4) is the deuterium labeled Valproic acid. Valproic acid (VPA; 2-Propylpentanoic Acid) is an HDAC inhibitor, with  $IC_{50}$  in the range of 0.5 and 2 mM, also inhibits HDAC1 (IC<sub>so</sub>, 400 μM), and induces proteasomal degradation of HDAC2.

Cat. No.: HY-10585S

Purity:

Clinical Data: No Development Reported

Size:

## Valproic acid-d4-1

(VPA-d4-1; 2-Propylpentanoic Acid-d4-1) Cat. No.: HY-10585S4

Valproic acid-d4-1 (VPA-d4-1) is the deuterium labeled Valproic acid. Valproic acid (VPA; 2-Propylpentanoic Acid) is an HDAC inhibitor, with IC<sub>so</sub> in the range of 0.5 and 2 mM, also inhibits HDAC1 (IC  $_{50^{\prime}}$  400  $\mu M$  ), and induces proteasomal degradation of HDAC2.

Purity: >98%

Clinical Data: No Development Reported Size: 1 mg, 5 mg

Valproic acid-d7 sodium (Sodium Valproate-d7 sodium)

Valproic acid-d7 (Sodium Valproate-d7) sodium is the deuterium labeled Valproic acid (sodium salt).

Cat. No.: HY-17362

Cat. No.: HY-10585AS

Purity: >98%

Clinical Data: No Development Reported

Size: 1 ma, 10 ma

#### Vancomycin hydrochloride

Vancomycin hydrochloride is an antibiotic for the treatment of bacterial infections. It acts by inhibiting the second stage of cell wall synthesis of susceptible bacteria. Vancomycin also alters the permeability of the cell membrane and selectively inhibits ribonucleic acid synthesis.

99.66% Purity: Clinical Data: Launched

10 mM × 1 mL, 250 mg, 1 g, 5 g

#### Valproic acid-d15

(VPA-d15; 2-Propylpentanoic Acid-d15)

Valproic acid-d15 is the deuterium labeled Valproic acid, Valproic acid (VPA: 2-Propylpentanoic Acid) is an HDAC inhibitor, with  $IC_{50}$  in the range of 0.5 and 2 mM, also inhibits **HDAC1** ( $IC_{50}$ , 400  $\mu$ M), and induces proteasomal degradation of HDAC2.

Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Valproic acid-d4 sodium

(VPA-d4 sodium; 2-Propylpentanoic Acid-d4 sodium)

Valproic acid-d4 (VPA-d4) sodium is the deuterium labeled Valproic acid. Valproic acid (VPA; 2-Propylpentanoic Acid) is an HDAC inhibitor, with  ${\rm IC}_{\rm 50}$  in the range of 0.5 and 2 mM, also inhibits HDAC1 (IC<sub>so</sub>, 400 μM), and induces proteasomal degradation of HDAC2.

**Purity:** 

Clinical Data: No Development Reported

1 mg, 5 mg

# Valproic acid-d6

(VPA-d6; 2-Propylpentanoic Acid-d6)

Valproic acid-d6 (VPA-d6) is the deuterium labeled Valproic acid. Valproic acid (VPA) 2-Propylpentanoic Acid) is an HDAC inhibitor, with IC<sub>so</sub> in the range of 0.5 and 2 mM, also inhibits HDAC1 (IC  $_{so'}$  400  $\mu M$  ), and induces proteasomal degradation of HDAC2.

Purity: 98.71%

Clinical Data: No Development Reported

Size: 5 mg, 10 mg

#### Vancomycin

Vancomycin is an antibiotic for the treatment of

bacterial infections.

96.66% Purity: Clinical Data: Launched

Size 25 mg, 50 mg, 100 mg, 1 g

# Vandetanib

(ZD6474)

Vandetanib (D6474) is a potent, orally active inhibitor of VEGFR2/KDR tyrosine kinase activity (IC<sub>so</sub>=40 nM). Vandetanib also has activity versus the tyrosine kinase activity of VEGFR3/FLT4  $(IC_{50}=110 \text{ nM})$  and EGFR/HER1  $(IC_{50}=500 \text{ nM})$ .

Purity: 99.89% Clinical Data: Launched

10 mM × 1 mL, 10 mg, 25 mg, 50 mg, 100 mg, 500 mg



Cat. No.: HY-10585S2









Cat. No.: HY-10585S3







Cat. No.: HY-10585S1







#### Vandetanib hydrochloride

(ZD6474 hydrochloride)

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).

nM) and EGFR/HER1 ( $IC_{50}$ =5 Purity: >98% Clinical Data: Launched Size: 1 mg, 5 mg



Cat. No.: HY-10260B

#### Vandetanib trifluoroacetate

(ZD6474 trifluoroacetate)

Vandetanib trifluoroacetate (D6474 trifluoroacetate) is a potent, orally active inhibitor of VEGFR2/KDR tyrosine kinase activity ( $IC_{so}$ =40 nM).

Purity: >98%
Clinical Data: Launched
Size: 1 mg, 5 mg



Cat. No.: HY-10260A

#### Vandetanib-d4

Cat. No.: HY-10260S1

Vandetanib-d4 (ZD6474-d4) is the deuterium labeled Vandetanib. Vandetanib (ZD6474) is a potent, orally active inhibitor of VEGFR2/KDR tyrosine kinase activity (IC $_{so}$ =40 nM).



**Purity:** > 98%

Clinical Data:

Size: 2.5 mg, 1 mg, 5 mg, 10 mg

# Vandetanib-d6

(ZD6474-d6)

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).



Cat. No.: HY-10260S

**Purity:** >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Veliparib

(ABT-888) Cat. No.: HY-10129

Veliparib (ABT-888) is a potent PARP inhibitor, inhibiting PARP1 and PARP2 with  $K_i$ s of 5.2 and 2.9 nM, respectively.



Purity: 99.78% Clinical Data: Phase 3

Size:  $10 \text{ mM} \times 1 \text{ mL}$ , 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

#### Veliparib dihydrochloride

(ABT-888 dihydrochloride)

Veliparib (dihydrochloride) is a potent inhibitor of PARP1 a nd PARP2 with K,s of 5.2 nM and 2.9 nM in cell-free assays, respectively.



Cat. No.: HY-10130

Purity: 99.96% Clinical Data: Phase 3

Size:  $10 \text{ mM} \times 1 \text{ mL}$ , 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

#### Vemurafenib

(PLX4032; RG7204; RO5185426) Cat. No.: HY-12057

Vemurafenib (PLX4032) is a first-in-class, selective, potent inhibitor of **B-RAF** kinase, with  $IC_{50}$ s of 31 and 48 nM for RAF $^{V600E}$  and c-RAF-1, respectively. Vemurafenib induces cell **autophagy**.



Purity: 99.83%
Clinical Data: Launched

Size:  $10 \text{ mM} \times 1 \text{ mL}, 10 \text{ mg}, 50 \text{ mg}, 100 \text{ mg}, 200 \text{ mg}, 500 \text{ mg}, 1 \text{ g}$ 

#### Vemurafenib-d5

Vemurafenib-d5 (PLX4032-d5) is the deuterium labeled Vemurafenib. Vemurafenib (PLX4032) is a first-in-class, selective, potent inhibitor of B-RAF kinase, with IC $_{50}$ s of 31 and 48 nM for RAF $^{V6006}$  and c-RAF-1, respectively. Vemurafenib induces cell **autophagy**.

Purity: >98%

Clinical Data:

Size: 1 mg, 5 mg, 10 mg, 25 mg, 50 mg



Cat. No.: HY-12057S

#### Vemurafenib-d7

(PLX4032-d7; RG7204-d7; RO5185426-d7) Cat. No.: HY-12057S1

Vemurafenib-d7 is deuterium labeled Vemurafenib. Vemurafenib (PLX4032) is a first-in-class, selective, potent inhibitor of B-RAF kinase, with IC50s of 31 and 48 nM for RAFV600E and c-RAF-1, respectively. Vemurafenib induces cell autophagy.



**Purity:** > 98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### Venetoclax

(ABT-199; GDC-0199)

Venetoclax (ABT-199; GDC-0199) is a highly potent, selective and orally bioavailable Bcl-2 inhibitor with a  $K_i$  of less than 0.01 nM. Venetoclax induces autophagy.



Cat. No.: HY-15531

Purity: 99.95% Clinical Data: Launched

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

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#### Venetoclax-d8

(ABT-199-d8; GDC-0199-d8) Cat. No.: HY-15531S

Venetoclax-d8 is deuterium labeled Venetoclax. Venetoclax (ABT-199: GDC-0199) is a highly potent. selective and orally bioavailable Bcl-2 inhibitor with a Ki of less than 0.01 nM. Venetoclax induces autophagy.



>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg



#### Verteporfin

(CL 318952) Cat. No.: HY-B0146

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

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Vinblastine sulfate

VER-155008

for Hsp70.

Purity:

Size:

(Vincaleukoblastine sulfate salt)

99 87%

Clinical Data: No Development Reported

Vinblastine sulfate is a cytotoxic alkaloid used against various cancer types. Vinblastine sulfate inhibits the formation of microtubule and suppresses nAChR with an  $IC_{50}$  of 8.9  $\mu$ M.

VER-155008 is an inhibitor of Hsp70, with IC<sub>so</sub>s

of 0.5  $\mu$ M, 2.6  $\mu$ M, and 2.6  $\mu$ M for Hsp70, Hsc70

and Grp7, respectively, and with a  $\boldsymbol{K}_{_{\!d}}$  of 0.3  $\mu M$ 



Cat. No.: HY-13780

Cat. No.: HY-10941

**Purity:** 99.04% Clinical Data: Launched

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

#### Vinorelbine ditartrate

(KW-2307; Nor-5'-anhydrovinblastine ditartrate) Cat. No.: HY-12053A

Vinorelbine (ditartrate) is an anti-mitotic agent which inhibits the proliferation of Hela cells with IC<sub>50</sub> of 1.25 nM.



98.08% Purity: Clinical Data: Launched

Size: 10 mM × 1 mL, 10 mg, 50 mg

#### Vinorelbine-d3 ditartrate (KW-2307-d3 ditartrate;

Nor-5'-anhydrovinblastine-d3 ditartrate) Cat. No.: HY-12053AS

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Vinorelbine-d3 (KW-2307-d3) ditartrate is the deuterium labeled Vinorelbine ditartrate. Vinorelbine (ditartrate) is an anti-mitotic agent which inhibits the proliferation of Hela cells with IC<sub>so</sub> of 1.25 nM.



>98% Purity:

Clinical Data: No Development Reported

Size 1 mg, 5 mg

#### Vismodegib

(GDC-0449) Cat. No.: HY-10440

Vismodegib (GDC-0449) is an orally active hedgehog pathway inhibitor with an IC<sub>50</sub> of 3 nM. Vismodegib also inhibits P-gp, ABCG2 with IC<sub>so</sub> values of 3.0 μM and 1.4 μM, respectively.



99.97% Purity: Clinical Data: Launched

Size 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg

#### Vistusertib (AZD2014)

Vistusertib (AZD2014) is an ATP competitive mTOR inhibitor with an  $IC_{50}$  of 2.81 nM. AZD2014 inhibits both mTORC1 and mTORC2 complexes.



Cat. No.: HY-15247

98.21% Purity: Clinical Data: Phase 2

Size 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

#### Vistusertib-d3

(AZD2014-d3) Cat. No.: HY-15247S

Vistusertib-d3 (AZD2014-d3) is the deuterium labeled Vistusertib. Vistusertib (AZD2014) is an ATP competitive mTOR inhibitor with an IC<sub>so</sub> of 2.81 nM. AZD2014 inhibits both mTORC1 and mTORC2 complexes.



Purity: >98%

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### VLX600

VLX600 is an iron-chelating inhibitor of oxidative phosphorylation (OXPHOS). VLX600 causes mitochondrial dysfunction and induces a strong shift to glycolysis. VLX600 displays selective

cytotoxic activity against malignant cell and induces autophagy. Anticancer activity.

Purity: 99.35%

Clinical Data: No Development Reported

5 mg, 10 mg, 25 mg, 50 mg, 100 mg, 250 mg

Cat. No.: HY-12406

#### VO-Ohpic trihydrate

Cat. No.: HY-13074

VO-Ohpic trihydrate is a highly potent inhibitor of PTEN with an IC<sub>50</sub> of 46±10 nM.

>98.0% Purity:

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg, 200 mg

#### Vorinostat

(SAHA; Suberoylanilide hydroxamic acid)

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<sub>50</sub> values of 10 nM and 20 nM for HDAC1 and HDAC3, respectively. Vorinostat induces cell apoptosis.

Cat. No.: HY-10221

Purity: 99 90% Clinical Data: Launched

10 mM × 1 mL, 250 mg, 500 mg, 1 g, 5 g

#### Vorinostat-d5

(SAHA-d5; Suberoylanilide hydroxamic acid-d5) Cat. No.: HY-115412

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<sub>so</sub> values of 10 nM and 20 nM for HDAC1 and HDAC3, respectively.

**Purity:** 

Clinical Data: No Development Reported

Size:

#### Vps34-IN-1

Vps34-IN-1 is an inhibitor of Vps34 extracted from patent WO2012085815A1, compound example 16a, with an IC<sub>so</sub> of 4 nM. Vps34-IN-1 modulates autophagy.



Cat. No.: HY-12795

**Purity:** 99.56%

Clinical Data: No Development Reported

10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

#### Vps34-IN-4

Cat. No.: HY-123058

Vps34-IN-4 (compound 19) is a potent, selective, and orally active inhibitor of VPS34. Vps34-IN-4 inhibits the autophagy in vivo. Autophagy is a dynamic process that regulates lysosomal-dependent degradation of cellular components.

>98% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

## Vps34-PIK-III

Cat. No.: HY-12794

Vps34-PIK-III is a potent and selective inhibitor of VPS34 with an IC<sub>50</sub> of 18 nM.

99.02% Purity:

Clinical Data: No Development Reported

10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 25 mg, 50 mg, 100 mg Size

#### VX-702

Cat. No.: HY-10401

VX-702 is a highly selective inhibitor of p38α MAPK, 14-fold higher potency against the p38α versus p38ß.

Purity: 99.44% Clinical Data: Phase 2

Size 10 mM × 1 mL, 10 mg, 50 mg, 100 mg, 200 mg

#### Wogonin

Wogonin is a naturally occurring mono-flavonoid, can inhibit the activity of CDK8 and Wnt, and exhibits anti-inflammatory and anti-tumor effects.

Cat. No.: HY-N0400

Purity: 99.98%

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

#### Wogonoside

Cat. No.: HY-N0399

Wogonoside, a flavonoid glycoside isolated from Huangqin, possesses anti-inflammatory effects. Wogonoside induces autophagy in breast cancer cells by regulating MAPK-mTOR pathway.

Purity: 99.92%

Clinical Data: No Development Reported Size: 5 mg, 10 mg, 20 mg

#### Wortmannin

(SL-2052; KY-12420)

Wortmannin (SL-2052; KY-12420) is a potent, selective and irreversible PI3K inhibitor with an IC<sub>so</sub> of 3 nM. Wortmannin also blocks autophagy formation, and potently inhibits Polo-like kinase 1 (PIK1) and PIk3 with  $IC_{50}$ s of 5.8 and 48 nM, respectively

Purity: 99.85%

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



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#### WYC-209

WYC-209, a synthetic retinoid, is a **retinoic acid receptor (RAR)** agonist. WYC-209 induces apoptosis primarily via the **caspase 3** pathway (IC $_{50}$ =0.19 $\mu$ M for inmalignant murine melanoma TRCs), and has long-term effects with little

toxicity.

Purity: 99.64%

Clinical Data: No Development Reported
Size: 5 mg, 10 mg, 50 mg, 100 mg



Cat. No.: HY-124136

## WYE-354

WYE-354 is an ATP-competitive mTOR inhibitor with an IC $_{50}$  of 5 nM. WYE-354 also inhibits PI3K $\alpha$  and PI3K $\gamma$  with IC $_{50}$ s of 1.89  $\mu$ M and 7.37  $\mu$ M, respectively. WYE-354 inhibits both mTORC1 and mTORC2. WYE-354 induces autophagy activation in vitro

Purity: 98.0%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 25 mg, 50 mg



Cat. No.: HY-12034

#### Xantocillin

(Xanthocillin X) Cat. No.: HY-122404

Xantocillin (Xanthocillin X) is a marine agent extracted from Penicillium commune, induces autophagy through inhibition of the MEK/ERK pathway.

**Purity:** >98%

Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg

#### XCT790

XCT-790 is a potent and selective inverse agonist for ERR $\alpha$  with an IC $_{50}$  value of 0.37  $\mu$ M. XCT-790 induces cell death in chemotherapeutic resistant cancer cells. XCT-790 (Compound 12) is inactive against ERR $\gamma$  and the estrogen receptors ER $\alpha$  and

ERβ.

**Purity:** 99.66%

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 10 mg, 50 mg



Cat. No.: HY-10426

XL388

Cat. No.: HY-13806

XL388 is a highly potent and ATP-competitive mTOR inhibitor with an  $\rm IC_{50}$  of 9.9 nM. XL388 simultaneously inhibits both mTORC1 and mTORC2.

**Purity:** 99.25%

Clinical Data: No Development Reported

Size:  $10 \text{ mM} \times 1 \text{ mL}$ , 5 mg, 10 mg, 50 mg, 100 mg

#### XRK3F2

Cat. No.: HY-112904

XRK3F2 is an inhibitor of **p62** (Sequestosome-1)-ZZ/ domain.



Purity: 98.85%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 5 mg, 10 mg, 50 mg, 100 mg

Xylitol

(Xylite) Cat. No.: HY-N0538

Xylitol is a chemical categorized as a polyalcohol or sugar alcohol. Target: Others Xylitol is a chemical categorized as a polyalcohol or sugar alcohol (alditol). Xylitol has the formula (CHOH)3(CH2OH)2 and is an achiral isomer of pentane-1,2,3,4,5-pentol.

HO OH OH

Purity: ≥98.0% Clinical Data: Launched

Size:  $10 \text{ mM} \times 1 \text{ mL}, 100 \text{ mg}$ 

#### Yangonin

Yangonin exhibits affinity for the human recombinant cannabinoid CB1 receptor with an IC  $_{so}$  and a  $\rm K_i$  of 1.79  $\mu M$  and 0.72  $\mu M$ ,

respectively.

**Purity:** 99.72%

Clinical Data: No Development Reported
Size: 10 mM × 1 mL, 5 mg, 10 mg

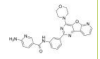


Cat. No.: HY-N0919

YM-201636

Cat. No.: HY-13228

YM-201636 is a potent and selective **PIKfyve** inhibitor with an IC $_{50}$  of 33 nM. YM-201636 also inhibits p110 $\alpha$  with an IC $_{50}$  of 3.3  $\mu$ M. YM-201636 inhibits **retroviral** replication.



**Purity:** 98.01%

Clinical Data: No Development Reported

Size: 10 mM × 1 mL, 2 mg, 5 mg, 10 mg, 50 mg, 100 mg

#### Zeaxanthin dipalmitate

(Physalien)

Zeaxanthin dipalmitate (Physalien) is a wolfberry-derived carotenoid, has anti-inflammatory and anti-oxidative stress

ty: >98%

Clinical Data: No Development Reported
Size: 5 mg, 10 mg, 25 mg



Cat. No.: HY-N9182

#### Zebularine

(NSC309132; 4-Deoxyuridine)

Cat. No.: HY-13420

Zebularine (NSC309132; 4-Deoxyuridine) is a DNA methyltransferase inhibitor. Zebularine also inhibits cytidine deaminase with a K, of 0.95 μΜ.

99 62% Purity:

Clinical Data: No Development Reported Size: 10 mM × 1 mL, 10 mg, 50 mg

#### Zingiberene

(α-Zingiberene; (-)-Zingiberene)

Zingiberene (α-Zingiberene) is a monocyclic sesquiterpene which is the predominant constituent of ginger with oil content (Zingiber officinale). Neuroprotective potential. Zingiberene triggers autophagy. Anticancer activity.



Cat. No.: HY-14618

97 67% Purity:

Clinical Data: No Development Reported

Size: 1 mg, 5 mg

#### **ZLN005**

Cat. No.: HY-17538

ZLN005 is a potent activator of peroxisome proliferator-activated receptor- $\gamma$  coactivator- $1\alpha$  $(PGC-1\alpha).$ 

**Purity:** 99 92%

Clinical Data: No Development Reported 10 mM × 1 mL, 10 mg, 50 mg

#### ZLN005-d4

Cat. No.: HY-17538S

ZLN005-d4 is deuterium labeled ZLN005. ZLN005 is a potent activator of peroxisome proliferator-activated receptor- $\gamma$  coactivator- $1\alpha$ 

 $(PGC-1\alpha)$ .

**Purity:** >98%

Clinical Data: No Development Reported

1 mg, 5 mg

#### Zoledronic Acid

(Zoledronate; CGP 42446; CGP42446A; ZOL 446) Cat. No.: HY-13777

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.



≥98.0% Purity: Clinical Data: Launched Size 50 mg, 100 mg

### Zoledronic acid monohydrate (Zoledronate monohydrate; CGP 42446 monohydrate; CGP42446A monohydrate; ...)

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.

H\_O\_H

Cat. No.: HY-50847

Cat. No.: HY-13777A

Purity: Clinical Data: Launched Size: 1 mg, 5 mg

>98%

#### **ZPCK**

(SL-01) Cat. No.: HY-100709

ZPCK is an oral active prodrug of gemcitabine that was designed for improved oral bioavailability.

Purity: >98%

Clinical Data: No Development Reported

Size 1 mg, 5 mg

#### ZSTK474

ZSTK474 is an ATP-competitive pan-class I PI3K inhibitor with IC<sub>so</sub>s of16 nM, 44 nM, 4.6 nM and 49 nM for PI3Kα, PI3Kβ, PI3Kδ and PI3Kγ,

respectively.

Purity: 99.71% Clinical Data: Phase 1

Size: 10 mg, 50 mg, 100 mg, 200 mg



ZX-29

Cat. No.: HY-135887

ZX-29 is a potent and selective ALK inhibitor with an IC<sub>50</sub> 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: 99.52%

Clinical Data: No Development Reported Size: 5 mg, 10 mg, 25 mg, 50 mg, 100 mg

#### α-Thujone

Cat. No.: HY-121618

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  $\alpha$ -Thujone is 21  $\mu$ M in suppressing the GABA-induced currents.

Purity: ≥95.0%

Clinical Data: No Development Reported

50 mg, 100 mg



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## β-Lapachone

(ARQ-501; NSC-26326)

Cat. No.: HY-13555

 $\beta$ -Lapachone (ARQ-501;NSC-26326) is a naturally occurring O-naphthoquinone, acts as a **topoisomerase I** inhibitor, and induces apoptosis by inhibiting cell cycle progression.

Purity: 99.85% Clinical Data: Phase 2

Size:  $10 \text{ mM} \times 1 \text{ mL}$ , 10 mg, 50 mg, 100 mg