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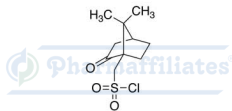
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Camphorquinone

Camphorquinone, also known as 2,3-bornanedione, is a photoinitiator used in curing dental composites. It is induced very slowly by camphorquinone.. Reference standards of Camphorquinone API, and its pharmacopeial, non pharmacopeial impurities, and stable isotopes are listed below



(1R)-(-)-10-Camphorsulfonyl Chloride

Catalogue No.: **PA 03 05580**

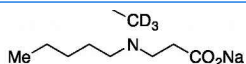
CAS : **39262-22-1**

Molecular Formula : **C₁₀H₁₅ClO₃S**

Molecular Weight : **250.74**

Ibandronic Acid

Ibandronic Acid is used prevention and treatment of osteoporosis and metastasis-associated skeletal fractures in people with cancer. It is also called bisphosphonate. Reference standards of Ibandronic Acid API, and its pharmacopeial, non pharmacopeial impurities, and stable isotopes are listed below.



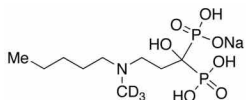
3-(N-Methyl-d3-N-pentylamino)propionic Acid Sodium Salt

Catalogue No.: **PA STI 064300**

CAS : **NA**

Molecular Formula : **C₉H₁₅D₃NNaO₂**

Molecular Weight : **198.25**



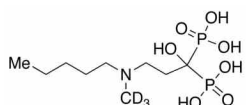
Ibandronic Acid-d3 Sodium Salt See I120003

Catalogue No.: **PA STI 052350**

CAS : **1329834-28-7**

Molecular Formula : **C₉H₁₅D₃NNaO₇P₂**

Molecular Weight : **344.23**



Ibandronic Acid-d3

Catalogue No.: **PA STI 052360**

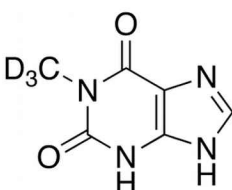
CAS : **1130899-41-0**

Molecular Formula : **C₉H₂₀D₃NO₇P₂**

Molecular Weight : **322.25**

Xanthine

Xanthine is used for their effects as mild stimulants and as bronchodilators, notably in the treatment of asthma or influenza symptoms. It is an intermediate in the degradation of adenosine monophosphate to uric acid, being formed by oxidation of hypoxanthine. Reference standards of Xanthine API, and its pharmacopeial, non pharmacopeial impurities, and stable isotopes are listed below.



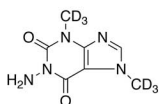
1-Methyl Xanthine-d3

Catalogue No.: **PA STI 064880**

CAS : **1216430-61-3**

Molecular Formula : **C₆H₃D₃N₄O₂**

Molecular Weight : **169.16**



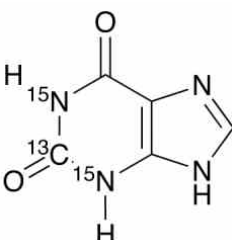
5-Amino-3,7-dimethyl Xanthine-d6

Catalogue No.: **PA STI 005960**

CAS : **1185126-58-2**

Molecular Formula : **C₇H₃D₆N₅O₂**

Molecular Weight : **201.22**



Xanthine-13C15N2

Catalogue No.: **PA STI 087470**

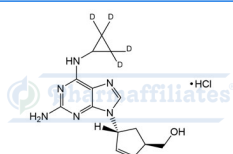
CAS : **1262670-81-4**

Molecular Formula : **C₄₁₃CH₄N₂₁₅N₂O₂**

Molecular Weight : **155.09**

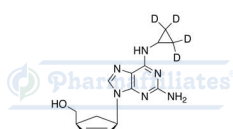
Abacavir

Abacavir is a chiral purine carbocyclic nucleoside, which is a prodrug of the guanosine analog (-) carbovir, Abacavir or ABC is an antiretroviral medication used to prevent and treat HIV/AIDS. Reference standards of Abacavir API, and its pharmacopeial, non pharmacopeial impurities, and stable isotopes are listed below.



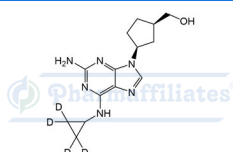
trans-Abacavir-d4 Hydrochloride

Catalogue No.: **PA STI 001020** CAS : **1346605-40-0** Molecular Formula : **C₁₄H₁₅D₄ClN₆O** Molecular Weight : **326.82**



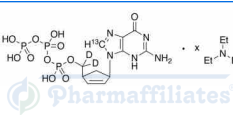
Abacavir-d4

Catalogue No.: **PA STI 001030** CAS : **1260619-56-4** Molecular Formula : **C₁₄H₁₄D₄N₆O** Molecular Weight : **290.36**



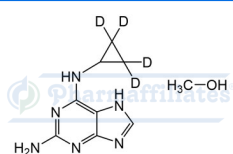
Dihydro Abacavir-D4

Catalogue No.: **PA STI 005680** CAS : **NA** Molecular Formula : **C₁₄H₁₆D₄N₆O** Molecular Weight : **292.37**



Carbovir-13C,d2 Triphosphate Triethylamine Salt

Catalogue No.: **PA STI 017680** CAS : **NA** Molecular Formula : **C₁₀₁₃CH₁₄D₂N₅O₁₁P₃•xC₆H₁₅N** Molecular Weight : **490.2**

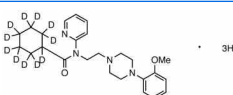


N6-Cyclopropyl-9H-purine-2,6-diamine-d4 Methanolate

Catalogue No.: **PA STI 023780** CAS : **NA** Molecular Formula : **C₉H₁₀D₄N₆O** Molecular Weight : **226.27**

Way 100635

WAY-100635 is a piperazine drug and research chemical widely used in scientific studies. It was originally believed to act as a selective 5-HT_{1A} receptor antagonist, but subsequent research showed that it also acts as potent full agonist at the D₄ receptor. It is sometimes referred to as a silent antagonist at the former receptor. It is closely related to WAY-100135. Reference standards of Way 100635 API, and its pharmacopeial, non pharmacopeial impurities, and stable isotopes are listed below.

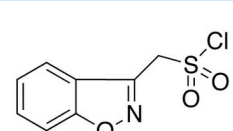


WAY 100635-d11 Hydrochloride

Catalogue No.: **PA STI 087440** CAS : **1329656-80-5** Molecular Formula : **C₂₅H₂₆D₁₁Cl₃N₄O₂** Molecular Weight : **543.01**

Zonisamide

Zonisamide is used to treat the symptoms of epilepsy and Parkinson's disease. It is a sulfonamide anticonvulsant. Zonisamide may be a carbonic anhydrase inhibitor although this is not one of the primary mechanisms of action. Reference standards of Zonisamide API, and its pharmacopeial, non pharmacopeial impurities, and stable isotopes are listed below.

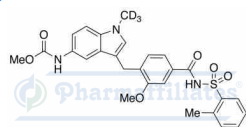


Benzo[d]isoxazol-3-yl-methanesulfonyl chloride

Catalogue No.: **PA 26 08520** CAS : **73101-65-2** Molecular Formula : **C₈H₆ClNO₃S** Molecular Weight : **231.66**

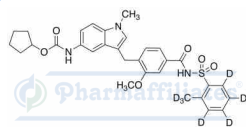
Zafirlukast

Zafirlukast is used to prevent asthma symptoms and to decrease the number of asthma attacks in people 5 and older. It is an orally administered leukotriene receptor antagonist. Reference standards of Zafirlukast API, and its pharmacopeial, non pharmacopeial impurities, and stable isotopes are listed below.



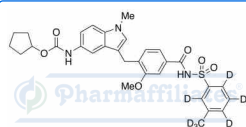
Decyclopentyl Zafirlukast-d3 Methyl Ester

Catalogue No.: **PA STI 024800** CAS : **1795011-90-3** Molecular Formula : **C₂₇H₂₄D₃N₃O₆S** Molecular Weight : **524.60**



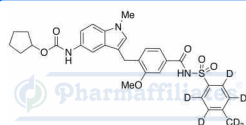
Zafirlukast-d7

Catalogue No.: **PA STI 087570** CAS : **1217174-18-9** Molecular Formula : **C₃₁H₂₆D₇N₃O₆S** Molecular Weight : **582.72**



Zafirlukast m-Tolyl Isomer-d7

Catalogue No.: **PA STI 087580** CAS : **1794760-52-3** Molecular Formula : **C₃₁H₂₆D₇N₃O₆S** Molecular Weight : **582.72**

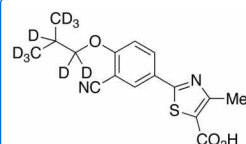


Zafirlukast p-Tolyl Isomer-d7

Catalogue No.: **PA STI 087590** CAS : **1794886-13-7** Molecular Formula : **C₃₁H₂₆D₇N₃O₆S** Molecular Weight : **582.72**

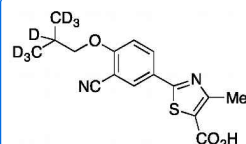
Febuxostat

Febuxostat is used to treat chronic gout and hyperuricemia. It is sold under the brand names Uloric among others, is a medication used long-term to treat gout due to high uric acid levels.. Reference standards of Febuxostat API, and its pharmacopeial, non pharmacopeial impurities, and stable isotopes are listed below



Febuxostat-d9

Catalogue No.: **PA STI 040630** CAS : **1246819-50-0** Molecular Formula : **C₁₆H₇D₉N₂O₃S** Molecular Weight : **325.43**

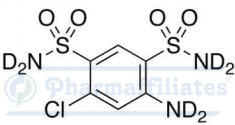
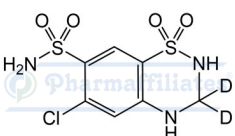
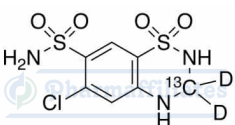
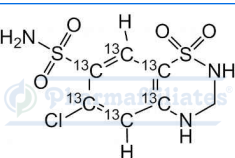
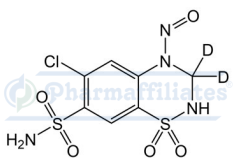
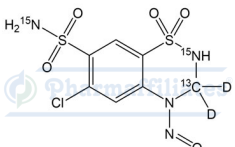


Febuxostat-d7

Catalogue No.: **PA STI 040640** CAS : **1285539-74-3** Molecular Formula : **C₁₆H₉D₇N₂O₃S** Molecular Weight : **323.42**

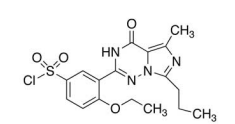
Hydrochlorothiazide

Hydrochlorothiazide is a drug that belongs to thiazide class of diuretics. It is used for the treatment of hypertension, congestive heart failure, symptomatic edema, diabetes insipidus, renal tubular acidosis. It is also used for treating kidney stones. Reference standards of Hydrochlorothiazide API, and its pharmacopeial, non pharmacopeial impurities, and stable isotopes are listed below.

	4-Amino-6-chloro-1,3-benzenedisulfonamide-d6	Catalogue No.: PA STI 005550	CAS : 1184995-49-0	Molecular Formula : C₆H₂D₆ClN₃O₄S₂	Molecular Weight : 291.77
	Hydrochlorothiazide-d2	Catalogue No.: PA STI 047000	CAS : 1219798-89-6	Molecular Formula : C₇H₆D₂ClN₃O₄S₂	Molecular Weight : 299.75
	Hydrochlorothiazide-13C,d2	Catalogue No.: PA STI 047010	CAS : 1190006-03-1	Molecular Formula : C₆₁₃CH₆D₂ClN₃O₄S₂	Molecular Weight : 300.75
	Hydrochlorothiazide-13C6	Catalogue No.: PA STI 047011	CAS : 1261396-79-5	Molecular Formula : C₁₃C₆H₈ClN₃O₄S₂	Molecular Weight : 303.70
	6-Chloro-4-nitroso-3,4-dihydro-2H-benzo[e][1,2,4]thiadiazine-7-sulfonamide-3,3-d2 1,1-dioxide	Catalogue No.: PA STI 089202	CAS : NA	Molecular Formula : C₇H₅D₂ClN₄O₅S₂	Molecular Weight : 328.74
	4-Nitroso Hydrochlorothiazide-13C,15N2,d2	Catalogue No.: PA STI 089441	CAS : NA	Molecular Formula : C₆₁₃CH₅D₂ClN₂₁₅N₂O₅S₂	Molecular Weight : 331.72

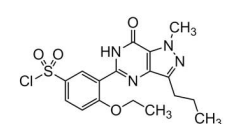
Vardenafil

Vardenafil is used to treat erectile dysfunction in men. It is an oral therapy for the treatment of erectile dysfunction. It is sold under the trade names Levitra, Staxyn, and Vivanza. Reference standards of Vardenafil API, and its pharmacopeial, non pharmacopeial impurities, and stable isotopes are listed below.

	4-Ethoxy-3-(5-methyl-4-oxo-7-propyl-3,4-dihydro-imidazo[5,1-f][1,2,4]-triazin-2-yl)benzene-sulfonyl Chloride	Catalogue No.: PA 22 14530	CAS : 224789-26-8	Molecular Formula : C₁₇H₁₉ClN₄O₄S	Molecular Weight : 410.88
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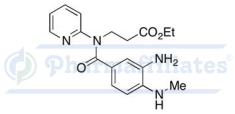
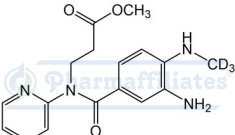
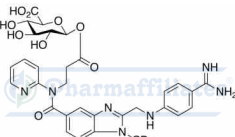
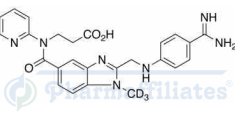
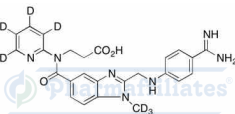
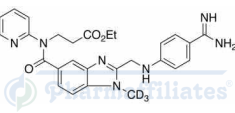
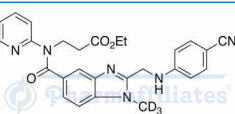
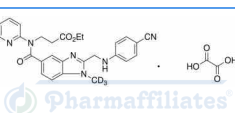
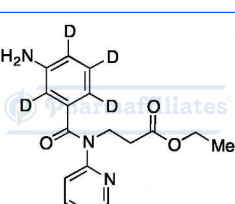
Sildenafil Citrate

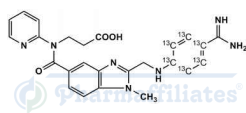
A citrate salt of Sildenafil sold under the brand name of Viagra. Sildenafil is used for the treatment of erectile dysfunction and pulmonary arterial hypertension. Sildenafil inhibits an enzyme cGMP-specific phosphodiesterase type 5 that promotes degradation of cGMP, which regulates blood flow in the penis. Reference standards of Sildenafil Citrate API, and its pharmacopeial, non pharmacopeial impurities, and stable isotopes are listed below.

	Sildenafil Chlorosulfonyl Impurity	Catalogue No.: PA 19 09570	CAS : 139756-22-2	Molecular Formula : C₁₇H₁₉ClN₄O₄S	Molecular Weight : 410.88
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Dabigatran

Dabigatran, sold under the brand name Pradaxa. It is used to prevent strokes in those with atrial fibrillation not caused by heart valve issues, as well as deep vein thrombosis and pulmonary embolism in persons who have been treated for 5–10 days with parenteral anticoagulant and to prevent deep vein thrombosis and pulmonary embolism in some circumstances.. Reference standards of Dabigatran API, and its pharmacopeial, non pharmacopeial impurities, and stable isotopes are listed below

	N-[3-Amino-4-(methylamino)benzoyl]-N-2-pyridinyl-β-alanine Ethyl Ester	Catalogue No.: PA 04 01650	CAS : 212322-56-0	Molecular Formula : C₁₈H₂₂N₄O₃	Molecular Weight : 342.39
	N-[3-Amino-4-(methylamino)benzoyl]-N-2-pyridinyl-β-alanine-d3 Methyl Ester	Catalogue No.: PA STI 006550	CAS : NA	Molecular Formula : C₁₇H₁₇D₃N₄O₃	Molecular Weight : 331.38
	Dabigatran Acyl-β-D-Glucuronide-d3	Catalogue No.: PA STI 024060	CAS : NA	Molecular Formula : C₃₁H₃₀D₃N₇O₉	Molecular Weight : 650.65
	Dabigatran-d3	Catalogue No.: PA STI 024070	CAS : 1246817-44-6	Molecular Formula : C₂₅H₂₂D₃N₇O₃	Molecular Weight : 474.53
	Dabigatran-d7	Catalogue No.: PA STI 024080	CAS : NA	Molecular Formula : C₂₅H₁₈D₇N₇O₃	Molecular Weight : 478.55
	Dabigatran-d3 Ethyl Ester Hydrochloride Salt	Catalogue No.: PA STI 024090	CAS : NA	Molecular Formula : C₂₇H₂₇D₃ClN₇O₃	Molecular Weight : 539.04
	Deacetamidine Cyano Dabigatran-d3 Ethyl Ester	Catalogue No.: PA STI 024470	CAS : 1346597-93-0	Molecular Formula : C₂₇H₂₃D₃N₆O₃	Molecular Weight : 485.55
	Deacetamidine Cyano Dabigatran-d3 Ethyl Ester Oxalate	Catalogue No.: PA STI 024480	CAS : 1794780-07-6	Molecular Formula : C₂₉H₂₅D₃N₆O₇	Molecular Weight : 575.59
	Ethyl 3-(3-Amino-N-(pyridin-2-yl)benzamido)propanoate-d4	Catalogue No.: PA STI 038730	CAS : NA	Molecular Formula : C₁₇H₁₅D₄N₃O₃	Molecular Weight : 317.38
Dabigatran-13C6					



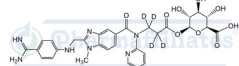
Catalogue No.: **PA STI 024081**

CAS : **1210608-88-0**

Molecular Formula : **C₁₉H₁₃C₆H₂₅N₇O₃**

Molecular Weight : **477.48**

Dabigatran β-acyl glucuronide D4



Catalogue No.: **PA STI 088975**

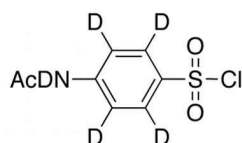
CAS : **NA**

Molecular Formula : **C₃₁H₂₉D₄N₇O₉**

Molecular Weight : **651.67**

Chitosan

Chitosan is used to treat obesity, high cholesterol, and Crohn's disease. It is a linear polysaccharide composed of randomly distributed β-(1→4)-linked D-glucosamine and N-acetyl-D-glucosamine. Reference standards of Chitosan API, and its pharmacopeial, non pharmacopeial impurities, and stable isotopes are listed below.



4-(Acetylamino)benzenesulfonyl-d5 Chloride

Catalogue No.: **PA STI 002300**

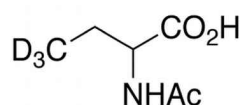
CAS : **1020718-84-6**

Molecular Formula : **C₈H₃D₅ClNO₃S**

Molecular Weight : **238.70**

Butanoic Acid

Butanoic Acid is an oily, colorless liquid that is soluble in water, ethanol, and ether. Butyric acid also known under the systematic name butanoic acid. Reference standards of Butanoic Acid API, and its pharmacopeial, non pharmacopeial impurities, and stable isotopes are listed below.



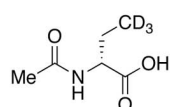
(+/-)-2-Acetylamino butanoic Acid-d3

Catalogue No.: **PA STI 002120**

CAS : **1219397-36-0**

Molecular Formula : **C₆H₈D₃NO₃**

Molecular Weight : **148.17**



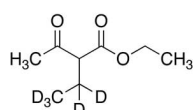
(2R)-2-(Acetylamino)butanoic Acid-d3

Catalogue No.: **PA STI 002150**

CAS : **2714420-85-4**

Molecular Formula : **C₆H₈D₃NO₃**

Molecular Weight : **148.18**



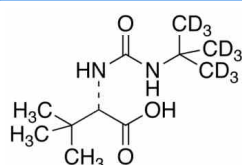
2-Acetylbutanoic-d5 Acid Ethyl Ester

Catalogue No.: **PA STI 002360**

CAS : **141327-44-8**

Molecular Formula : **C₈H₉D₅O₃**

Molecular Weight : **163.23**



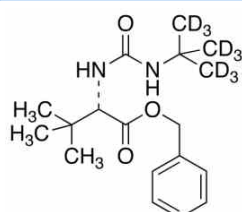
N-tert-Butylcarbamoyl-L-tert-leucine-d9

Catalogue No.: **PA STI 016070**

CAS : **NA**

Molecular Formula : **C₁₁H₁₃D₉N₂O₃**

Molecular Weight : **239.36**



N-tert-Butylcarbamoyl-L-tert-leucine-d9 Benzyl Ester

Catalogue No.: **PA STI 016080**

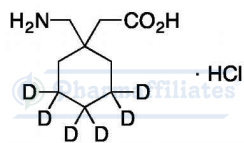
CAS : **1795786-80-9**

Molecular Formula : **C₁₈H₁₉D₉N₂O₃**

Molecular Weight : **329.48**

Gabapentin

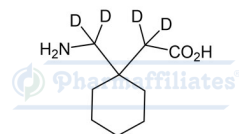
An anti-epileptic and anticonvulsant drug, Gabapentin is used to treat epilepsy, neuropathic pain, hot flashes and other anxiety disorders. It possibly inhibits neurotransmitter γ -aminobutyric acid (GABA). Reference standards of Gabapentin API, and its pharmacopeial, non pharmacopeial impurities, and stable isotopes are listed below.



Gabapentin-d6 Hydrochloride

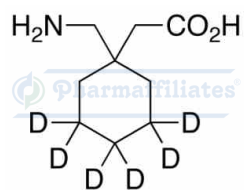
Catalogue No.: **PA STI 043690** CAS : **1432061-73-8** Molecular Formula : **C₉H₁₂D₆ClNO₂** Molecular Weight : **213.73**

SIMILAR PRODUCTS



Gabapentin-d4

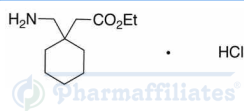
Catalogue No.: **PA STI 043700** CAS : **1185039-20-6** Molecular Formula : **C₉H₁₃D₄NO₂** Molecular Weight : **175.26**



Gabapentin-d6

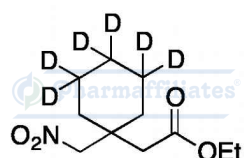
Catalogue No.: **PA STI 043710** CAS : **1346600-67-6** Molecular Formula : **C₉H₁₁D₆NO₂** Molecular Weight : **177.27**

SIMILAR PRODUCTS



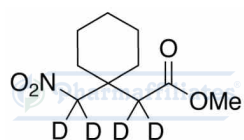
Gabapentin Ethyl Ester Hydrochloride

Catalogue No.: **PA STI 043720** CAS : **60175-04-4** Molecular Formula : **C₁₁H₂₁NO₂** Molecular Weight : **199.26**



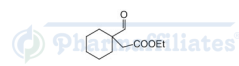
1-(Nitromethyl)cyclohexane-d6-acetic Acid Ethyl Ester

Catalogue No.: **PA STI 068190** CAS : **NA** Molecular Formula : **C₁₁H₁₃D₆NO₄** Molecular Weight : **235.31**



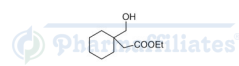
1-(Nitromethyl)cyclohexaneacetic Acid-d4 Methyl Ester

Catalogue No.: **PA STI 068200** CAS : **1246815-71-3** Molecular Formula : **C₁₀H₁₃D₄NO₄** Molecular Weight : **219.27**



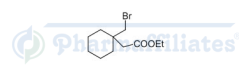
Ethyl (1-formylcyclohexyl)acetate,

Catalogue No.: **PAI 07 007020** CAS : **460711-33-5** Molecular Formula : **C₁₁H₁₈O₃** Molecular Weight : **198.26**



Ethyl 2-(1-(hydroxymethyl)cyclohexyl)acetate,

Catalogue No.: **PAI 07 007030** CAS : **NA** Molecular Formula : **C₁₁H₂₀O₃** Molecular Weight : **200.28**



Ethyl 2-(1-(bromomethyl)cyclohexyl)acetate,

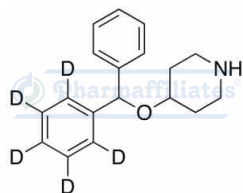
Catalogue No.: **PAI 07 007040** CAS : **NA** Molecular Formula : **C₁₁H₁₉BrO₂** Molecular Weight : **263.18**

Gabapentin-d4

Catalogue No.: **PA STI 088406** CAS : **1407511-35-6** Molecular Formula : **C₉H₁₃D₄NO₂** Molecular Weight : **175.26**

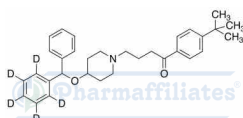
Ebastine

Ebastine is a second generation H1 antihistamine drug, which does not cross the blood brain barrier and causes low drowsiness. It is used for the treatment of rhinitis and urticaria. Reference standards of Ebastine API, and its pharmacopeial, non pharmacopeial impurities, and stable isotopes are listed below.



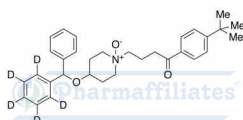
Desalkyl Ebastine-d5

Catalogue No.: **PA STI 027090** CAS : **1189424-36-9** Molecular Formula : **C₁₈H₁₆D₅NO** Molecular Weight : **272.40**



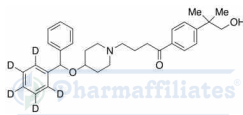
Ebastine-d5

Catalogue No.: **PA STI 036720** CAS : **1216953-13-7** Molecular Formula : **C₃₂H₃₄D₅NO₂** Molecular Weight : **474.69**



Ebastine-d5 N-Oxide

Catalogue No.: **PA STI 036730** CAS : **NA** Molecular Formula : **C₃₂H₃₄D₅NO₃** Molecular Weight : **490.69**

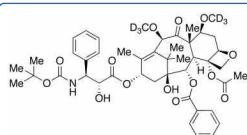


Hydroxy Ebastine-d5

Catalogue No.: **PA STI 048260** CAS : **1189954-07-1** Molecular Formula : **C₃₂H₃₄D₅NO₃** Molecular Weight : **490.69**

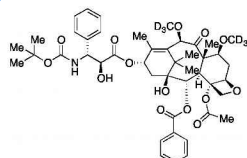
Cabazitaxel

Cabazitaxel was developed by Sanofi-Aventis and was approved by the U.S. FDA for the treatment of hormone-refractory prostate cancer on June 17, 2010. It is a semi-synthetic derivative of a natural taxoid. It is a microtubule inhibitor, and the fourth taxane to be approved as a cancer therapy.. Reference standards of Cabazitaxel API, and its pharmacopeial, non pharmacopeial impurities, and stable isotopes are listed below



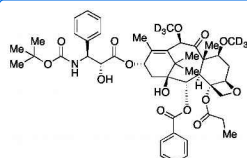
Cabazitaxel-d6

Catalogue No.: **PA STI 016870** CAS : **1383561-29-2** Molecular Formula : **C₄₅H₅₁D₆NO₁₄** Molecular Weight : **841.97**



(2'S, 3'R)-Cabazitaxel-D6

Catalogue No.: **PA STI 016880** CAS : **NA** Molecular Formula : **C₄₅H₅₁D₆NO₁₄** Molecular Weight : **841.97**



4-Deacetyl-4-propionyl Cabazitaxel-D6

Catalogue No.: **PA STI 024550** CAS : **NA** Molecular Formula : **C₄₆H₅₃D₆NO₁₄** Molecular Weight : **856.00**

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Россия +7(495)268-04-70

Казахстан +7(7172)727-132

Киргизия +996(312)96-26-47

эл.почта: peb@nt-rt.ru || сайт: <https://pharmaffiliates.nt-rt.ru/>