MCE RedChemExpress

Fenofibrate-d4

 Cat. No.:
 HY-17356S1

 CAS No.:
 1092484-57-5

 Molecular Formula:
 C₂₀H₁₇D₄ClO₄

Molecular Weight: 364.86

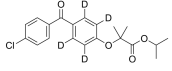
Target: PPAR; Autophagy; Cytochrome P450

Pathway: Cell Cycle/DNA Damage; Vitamin D Related/Nuclear Receptor; Autophagy; Metabolic

Enzyme/Protease

Storage: Please store the product under the recommended conditions in the Certificate of

Analysis.



BIOLOGICAL ACTIVITY

Description	Fenofibrate- d_4 is the deuterium labeled Fenofibrate[1]. Fenofibrate is a selective PPAR α agonist with an EC50 of 30 μ M. Fenofibrate also inhibits human cytochrome P450 isoforms, with IC50s of 0.2, 0.7, 9.7, 4.8 and 142.1 μ M for CYP2C19, CYP2B6, CYP2C9, CYP2C8, and CYP3A4, respectively[2][3].
In Vitro	Stable heavy isotopes of hydrogen, carbon, and other elements have been incorporated into drug molecules, largely as tracers for quantitation during the drug development process. Deuteration has gained attention because of its potential to affect the pharmacokinetic and metabolic profiles of drugs ^[1] . MCE has not independently confirmed the accuracy of these methods. They are for reference only.

REFERENCES

[1]. Russak EM, et al. Impact of Deuterium Substitution on the Pharmacokinetics of Pharmaceuticals. Ann Pharmacother. 2019 Feb;53(2):211-216.

[2]. Schelleman H, et al. Pharmacoepidemiologic and in vitro evaluation of potential drug-drug interactions of sulfonylureas with fibrates and statins. Br J Clin Pharmacol. 2014 Sep;78(3):639-48.

[3]. Gong Y, et al. Fenofibrate Inhibits Cytochrome P450 Epoxygenase 2C Activity to Suppress Pathological Ocular Angiogenesis. EBioMedicine. 2016 Sep 30. pii: S2352-3964(16)30448-0.

Caution: Product has not been fully validated for medical applications. For research use only.

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