

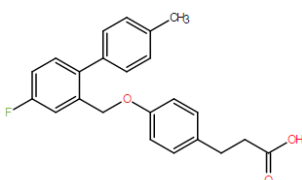
Certificate of Analysis

Target	生物活性小分子>>GPCR>>Free Fatty Acid Receptor (FFAR)
Cat.No	DC9981
Name	TUG891

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Chemical Properties

CAS	1374516-07-0
Formula	C ₂₃ H ₂₁ FO ₃
MW	364.41
Storage	2 years -20°C Powder, 2 weeks 4°C in DMSO, 6 months -80°C in DMSO
Structure	 <p>The chemical structure of TUG891 is a complex molecule. It features a central benzene ring with a fluorine atom at the 4-position and a methylene group at the 2-position. This methylene group is connected via an ether oxygen to another benzene ring, which has a methyl group at the 4-position. This second benzene ring is further connected via a propyl chain to a carboxylic acid group (-COOH).</p>
Purity	>98%

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www.dcchemicals.com