

产品名称: Darifenacin HBr

产品别名: Darifenacin hydrobromide; 氢溴酸达非那新

生物活性:

Description	Darifenacin hydrobromide (UK-88525 hydrobromide) is a selective M3 muscarinic receptor antagonist with pKi of 8.9. IC50 value: 8.9 (pKi) [1] Target: M3 receptor in vitro: Darifenacin exerts non-parallel rightward displacement of the agonist curve and also significant depression of the maximum response (+)-cis-Dioxolane produced concentration-dependent contraction of the isolated bladder of rat [1]. Darifenacin produces a concentration dependent increase in R123 (P-gp probe) accumulation in MDCK cells. Darifenacin stimulates ATPase activity in P-gp membrane in a clear concentration dependent response manner with an estimated ED50 value of 1.6 μ M. Darifenacin (100 nM) shows a significantly greater permeability for darifenacin in the basolateral to apical direction resulting in an efflux ratio in BBMEC monolayers of approximately 2.6 [2]. in vivo: Darifenacin produces dose-dependent inhibition of amplitude of volume-induced bladder contractions(VIBCAMP), producing 35% inhibition at dose of 283.3 nmol/kg and maximal inhibition of approximately 50–55% [1]. Darifenacin (0.1 mg/kg i.v.) reduces bladder afferent activity in both A δ and C fibers in female Sprague-Dawley rats, the decrease in afferent spikes in C fibers may be more pronounced than that in A δ fibers [3].																				
In Vitro: DMSO : 33.33 mg/mL (65.68 mM; Need ultrasonic) H ₂ O : < 0.1 mg/mL (insoluble)	<table border="1"><thead><tr><th rowspan="2"></th><th>Solvent / Mass Concentration</th><th>1 mg</th><th>5 mg</th><th>10 mg</th></tr></thead><tbody><tr><td>Preparing Stock Solutions</td><td>1 mM</td><td>1.9706 mL</td><td>9.8530 mL</td><td>19.7060 mL</td></tr><tr><td></td><td>5 mM</td><td>0.3941 mL</td><td>1.9706 mL</td><td>3.9412 mL</td></tr><tr><td></td><td>10 mM</td><td>0.1971 mL</td><td>0.9853 mL</td><td>1.9706 mL</td></tr></tbody></table>		Solvent / Mass Concentration	1 mg	5 mg	10 mg	Preparing Stock Solutions	1 mM	1.9706 mL	9.8530 mL	19.7060 mL		5 mM	0.3941 mL	1.9706 mL	3.9412 mL		10 mM	0.1971 mL	0.9853 mL	1.9706 mL
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Solvent&Solubility *请根据产品在不同溶剂中的溶解度选择合适的溶剂配制储备液 一旦配成溶液, 请分装保存, 避免反复冻融造成的产品失效。 储备液的保存方式和期限 -80°C, 6 months; -20°C, 1 month。 -80°C 储存时, 请在 6 个月内使用, -20°C 储存时, 请在 1 个月内使用。 In Vivo: 请根据您的实验动物和给药方式选择适当的溶解方案。以下溶解方案都请先按照 In Vitro 方式配制澄清的储备液, 再依次添加助溶剂: ——为保证实验结果的可靠性, 澄清的储备液可以根据储存条件, 适当保存; 体内实验的工作液, 建议您现用现配, 当天使用; 以下溶剂前显示的百分比是指该溶剂在您配制终溶液中的体积占比; 如在配制过程中出现沉淀、析出现象, 可以通过加热和/或超声的方式助溶 1.请依序添加每种溶剂: 10% DMSO→40% PEG300 →5% Tween-80 → 45% saline Solubility: ≥ 2.5 mg/mL (4.93 mM); Clear solution 此方案可获得 ≥ 2.5 mg/mL (4.93 mM, 饱和度未知) 的澄清溶液。 以 1 mL 工作液为例, 取 100 μ L 25.0 mg/mL 的澄清 DMSO 储备液加到 400 μ L PEG300 中, 混合均匀 向上述体系中加入 50 μ L Tween-80, 混合均匀; 然后继续加入 450 μ L 生理盐水定容至 1 mL。 2.请依序添加每种溶剂: 10% DMSO→ 90% (20% SBE-β-CD in saline) Solubility: ≥ 2.5 mg/mL (4.93 mM); Clear solution 此方案可获得 ≥ 2.5 mg/mL (4.93 mM, 饱和度未知) 的澄清溶液。																					

	<p>以 1 mL 工作液为例，取 100 μL 25.0 mg/mL 的澄清 DMSO 储备液加到 900 μL 20% 的 SBE-β-CD 生理盐水水溶液中，混合均匀。</p> <p>3.请依序添加每种溶剂： 10% DMSO → 90% corn oil Solubility: \geq 2.5 mg/mL (4.93 mM); Clear solution</p> <p>此方案可获得 \geq 2.5 mg/mL (4.93 mM, 饱和度未知) 的澄清溶液，此方案不适用于实验周期在半个月以上的实验。</p> <p>以 1 mL 工作液为例，取 100 μL 25.0 mg/mL 的澄清 DMSO 储备液加到 900 μL 玉米油中，混合均匀。</p>
References	<p>[1]. Hegde SS, et al. Functional role of M2 and M3 muscarinic receptors in the urinary bladder of rats <i>in vitro</i> and <i>in vivo</i>. <i>Br J Pharmacol</i>, 1997, 120(8), 1409-1418.</p> <p>[2]. Miller DW, et al. Evaluation of drug efflux transporter liabilities of darifenacin in cell culture models of the blood-brain and blood-ocular barriers. <i>Neurourol Urodyn</i>, 2011, 30(8), 1633-1638.</p> <p>[3]. Iijima K, et al. Effects of the M3 receptor selective muscarinic antagonist darifenacin on bladder afferent activity of the rat pelvic nerve. <i>Eur Urol</i>, 2007, 52(3), 842-847.</p>



源叶生物