

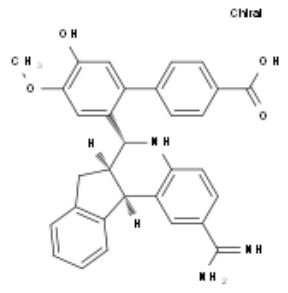
## Certificate of Analysis

Target	Products>>Novel inhibitors
Cat.No	DC72709
Name	BMS-593214

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## Chemical Properties

CAS	1004551-40-9
Formula	C <sub>31</sub> H <sub>27</sub> N <sub>3</sub> O <sub>4</sub>
MW	505.56
Storage	2 years -20°C Powder, 2 weeks 4°C in DMSO, 6 months -80°C in DMSO
Structure	 <p>The chemical structure shows a central chiral center (labeled 'Chiral') bonded to a 3,4-dihydroquinoline ring system, a 3,4-dihydroisoquinoline ring system, and a 2,4,6-trihydroxyphenyl group. The 2,4,6-trihydroxyphenyl group is further substituted with a methyl group and a carboxylic acid group. The 3,4-dihydroisoquinoline ring system is substituted with an amino group and a primary amide group.</p>
Purity	>98%

**Website:**  
www.dchemicals.com