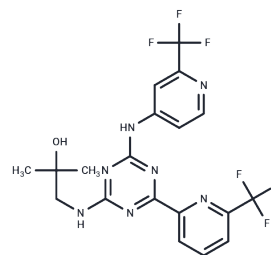


Enasidenib

Chemical Properties

CAS No. :	1446502-11-9
Formula:	C ₁₉ H ₁₇ F ₆ N ₇ O
Molecular Weight:	473.38
Appearance:	no data available
Storage:	Powder: -20°C for 3 years In solvent: -80°C for 1 year



Biological Description

Description	Enasidenib (AG-221) is an orally available inhibitor of specific mutant forms of the mitochondrial enzyme isocitrate dehydrogenase type 2 (IDH2), with potential antineoplastic activity.
Targets(IC50)	Dehydrogenase, Isocitrate Dehydrogenase (IDH)
In vitro	The compound has been demonstrated to reduce 2-HG levels by >90% and reverse histone and deoxyribonucleic acid (DNA) hypermethylation in vitro, and to induce differentiation in leukemia cell models[2].
In vivo	AG-221 is able to potently reduce 2HG found in the bone marrow, plasma and urine of engrafted mice. Treatment also induced a dose dependent, statistically significant, survival benefit. A proliferative burst of the human specific CD45+ blast cells is followed by cellular differentiation as measured by the expression of CD11b, CD14 and CD15 and cell morphology after AG-221 treatment[2]. AG-221 treatment also restores megakaryocyte-erythroid progenitor (MEP) differentiation that is suppressed by mutant IDH2 expression and reverses the effects of mutant IDH2 on DNA methylation in mutant stem/progenitor cells. Clinical trials combining IDH2 inhibitors with other targeted AML therapies are warranted in order to increase therapeutic efficacy[1].
Kinase Assay	Untranslated region-mediated luciferase reporter expression: HEK293 cells are transfected with a GEMS reporter vector that contains the luciferase open-reading frame flanked by and under post-transcriptional control of the BMI-1 5' and 3' UTRs. The resulting stable cells (F8) are treated with PTC-209 or vehicle control overnight, and then luciferase reporter activity is determined using Bright-Glo assays. The assays are run in triplicate for each point, and the percentage of inhibition was calculated against vehicle control.

Solubility Information

Solubility	H ₂ O: < 1 mg/mL (insoluble or slightly soluble), Ethanol: 100 mg/mL (211.25 mM), DMSO: 50 mg/mL (105.62 mM), < 1 mg/ml refers to the product slightly soluble or insoluble)
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A DRUG SCREENING EXPERT

Preparing Stock Solutions

	1mg	5mg	10mg
1 mM	2.1125 mL	10.5623 mL	21.1247 mL
5 mM	0.4225 mL	2.1125 mL	4.2249 mL
10 mM	0.2112 mL	1.0562 mL	2.1125 mL
50 mM	0.0422 mL	0.2112 mL	0.4225 mL

Please select the appropriate solvent to prepare the stock solution, according to the solubility of the product in different solvents. Please use it as soon as possible.

Reference

Alan H. Shih, et al. Blood. 2014, 124:437.

Kate Ellwood-Yen, et al. AACR. 2014, 74(19 Sup.):3116.

Inhibitor · Natural Compounds · Compound Libraries · Recombinant Proteins

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