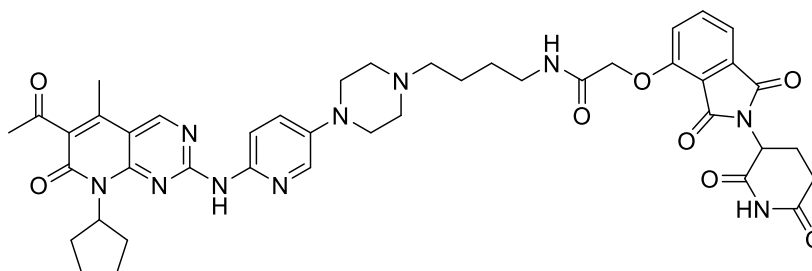
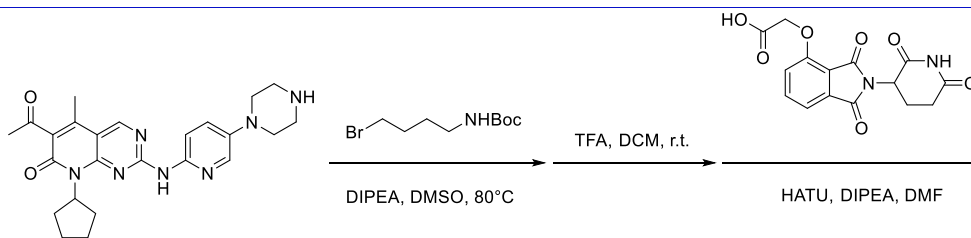


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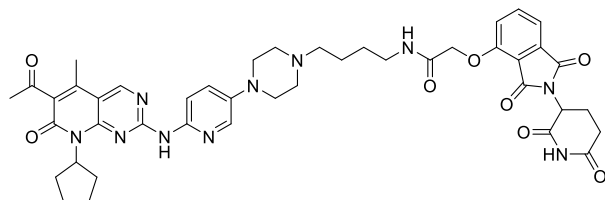


Chemical Formula: C₄₃H₄₈N₁₀O₈
Molecular Weight: 832.9190

Category	Parameter	Description
Compound	Name	BSJ-03-204
	Citation	Angew. Chem. Int. Edit., 2019, 58, 6321-6326. (https://onlinelibrary.wiley.com/doi/abs/10.1002/anie.201901336)
	Chemical descriptors	CC(C1=C(C2=CN=C(N=C2N(C1=O)C3CCCC3)NC4=CC=C(N5CCN(CC5)CCCCNC(COC6=CC=CC7=C6N(C7=O)C8CCC(NC8=O)=O)=O)=O)C=N4)C)=O
	Chemical name	N-(4-(4-(6-((6-acetyl-8-cyclopentyl-5-methyl-7-oxo-7,8-dihydropyrido[2,3-d]pyrimidin-2-yl)amino)pyridin-3-yl)piperazin-1-yl)butyl)-2-((2-(2,6-dioxopiperidin-3-yl)-1,3-dioxoisindolin-4-yl)oxy)acetamide
	Entries in chemical databases	
	Availability	
<i>In vitro</i> profiling	Papers that use the compounds	
	Additional comments	
	Target (potency)	CDK4/cyclin D1: 26.9 nM
	Target (potency)	CDK6/cyclin D1: 10.4 nM
	Selectivity	
	Potential reactivity	
Cellular profiling	SAR	
	Mechanism of inhibition	Protein degradation by recruiting Cereblon
	Structure of target-probe complex	
	Additional comments	
	Validation of cellular target	Dose-dependent degradation of CDK4 and CDK6 with maximum degradation at 0.25uM in Jurkat cells.
	Validation of cellular specificity	Selective degradation of CDK4 and CDK6 in Molt4 cells by proteomics analysis at 5h with 250 nM treatment.
Additional comments		
Pharmacodynamics		
Pharmacokinetics		



Synthetic scheme



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