



PDD 00017273

Catalog No: tcsc0028457

Av	ailable Sizes
Size: 5m	g
Size: 10r	ng
Size: 25r	ng
Sp	ecifications
CAS No: 1945950-	21-9
Formula	
Pathway Others	:
Target: Others	
Purity / 0 >98%	Grade:
Solubilit 10 mM in	
Observe 514.62	d Molecular Weight:

Product Description

PDD 00017273 is a potent inhibitor of Poly(ADP-ribose) Glycohydrolase (**PARG**), with an $\mathbf{IC}_{\mathbf{50}}$ of 26 nM, and a $\mathbf{K}_{\mathbf{D}}$ of 1.45 nM.

IC50 & Target: IC50: 26 nM (PARG)^[1]





KD: 1.45 nM (PARG)^[1]

In Vitro: PDD 00017273 is a potent inhibitor of PARG, with an IC $_{50}$ of 26 nM, and a K $_{D}$ of 1.45 nM. PDD 00017273 (10 μ M) does not inhibit five common Cytochrome P450 enzymes. PDD 00017273 (30 μ M) modestly increasess phosphorylated H2AX (γ H2AX) intensity, PDD 00017273 also decreases in NAD/H through PARG inhibition after DNA damage. PDD 00017273 suppresses the ZR-75-1 cells carring BRCA1 and BRCA2 wild type, and exhibits less potent activities against MDA-MB-436 cells carry the 5396 + 1G>A mutation in BRCA1 $^{[1]}$. PDD 00017273 (0.3 μ M) inhibits degradation of PAR polymers in MCF7 cells. PDD 00017273 (0.3 μ M) also reduces the viability of BRCA1, BRCA2, PALB2, FAM175A, and BARD1 depleted cells. PDD 00017273 stalls replication forks and induces DNA damage that requires homologous recombination (HR) for repair $^{[2]}$.

All products are for RESEARCH USE ONLY. Not for diagnostic & therapeutic purposes!