## **Product** Data Sheet

## PLX647 dihydrochloride

**Cat. No.:** HY-13838A **CAS No.:** 1779796-38-1

Molecular Formula: C<sub>21</sub>H<sub>19</sub>Cl<sub>2</sub>F<sub>3</sub>N<sub>4</sub>

Molecular Weight: 455.3

Target: c-Fms; c-Kit

Pathway: Protein Tyrosine Kinase/RTK

Storage: Please store the product under the recommended conditions in the Certificate of

Analysis.

## **BIOLOGICAL ACTIVITY**

**Description** PLX647 dihydrochloride is an orally active, highly specific dual FMS and KIT kinase inhibitor, with IC<sub>50</sub>s of 28 and 16 nM,

 $reapectively.\ PLX647\ dihydrochloride\ shows\ selectivity\ for\ FMS\ and\ KIT\ over\ a\ panel\ of\ 400\ kinases\ at\ a\ concentration\ of\ 1\ \mu$ 

M except FLT3 and KDR (IC<sub>50</sub>s=91 and 130 nM, respectively) $^{[1]}$ .

In Vitro In vitro, PLX647 dihydrochloride potently inhibits proliferation of BCR-FMS cells, with an IC<sub>50</sub> of 92 nM. A corresponding

Ba/F3 cell line expressing BCR-KIT is also quite sensitive to PLX647 dihydrochloride, with an IC $_{50}$  of 180 nM. PLX647 dihydrochloride also inhibits endogenous FMS and KIT, as demonstrated by inhibition of the ligand-dependent cell lines M-

NFS-60 (IC<sub>50</sub>=380 nM) and M-07e (IC<sub>50</sub>=230 nM), which express FMS and KIT, respectively<sup>[1]</sup>.

PLX647 dihydrochloride potently inhibits the growth of FLT3-ITD-expressing MV4-11 cells (IC $_{50}$ =110 nM). PLX647 dihydrochloride displays minimal inhibition of the proliferation of Ba/F3 cells expressing BCR-KDR (IC $_{50}$ =5  $\mu$ M). PLX647

dihydrochloride inhibits osteoclast differentiation with an IC  $_{50}$  of 0.17  $\mu M^{[1]}$ .

MCE has not independently confirmed the accuracy of these methods. They are for reference only.

In Vivo PLX647 dihydrochloride (40 mg/kg; p.o.; twice daily for 7 days) reduces macrophage accumulation in UUO kidney and blood monocytes<sup>[1]</sup>.

PLX647 dihydrochloride (40 mg/kg; p.o.; male Swiss Webster mice) reduces LPS-induced TNF-α and IL-6 release<sup>[1]</sup>.

PLX647 dihydrochloride (20-80 mg/kg; p.o.; daily or twice daily from 27-41 days) shows effects on collagen-induced arthritis

PLX647 dihydrochloride (30 mg/kg) results in significant inhibition of TRAP5b immunostaining and bone osteolysis. PLX647 dihydrochloride (30 mg/kg BID) is able to prevent bone damage by the tumor cells<sup>[1]</sup>.

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Animal Model:	Male C57BL/6 mice (mouse unilateral ureter obstruction model) <sup>[1]</sup>	
Dosage:	40 mg/kg	
Administration:	P.o.; twice daily for 7 days	
Result:	Resulted in reduction in the levels of F4/80+ macrophages by 77%.	
Animal Model:	7-9 wk old Male DBA/1J mice (Mouse collagen-induced arthritis model) $^{[1]}$	

Dosage:	20 mg/kg, 80 mg/kg
Administration:	P.o.; daily (20 mg/kg) from 27-41 days, twice daily (80 mg/kg) from 27-41 days
Result:	20 mg/kg PLX647 had no initial effect on the development of severe arthritis. However, starting on day 33, no further development of disease severity was recorded, and a 30% inhibition of the macroscopic signs of arthritis was evident in clinical score on day 41. Mic treated with 80 mg/kg BID PLX647 initially shows delayed development of severe arthritic signs. Starting on day 33, the signs of arthritis began to decrease in this treatment group, reaching a maximum reversal of 76% on day 41.

## **REFERENCES**

[1]. Zhang C, et al. Design and pharmacology of a highly specific dual FMS and KIT kinase inhibitor. Proc Natl Acad Sci U S A. 2013 Apr 2;110(14):5689-94.

 $\label{lem:caution:Product} \textbf{Caution: Product has not been fully validated for medical applications. For research use only.}$ 

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