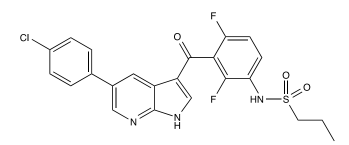


## Catalog # 10-2928 PLX-4032

918504-65-1

N-[3-[[5-(4-chlorophenyl)-1H-pyrrolo[2,3-b]pyridin-3-yl]carbonyl]-2,4-difluorophenyl]-1-propanesulfonamide; Vemurafenib; R7204; RG-7204; Ro 51-85426

Lot # X107376



An ATP-competitive inhibitor of mutant V600E/F and wild type B-Raf, IC<sub>50</sub>s=31 and 100 nM respectively.<sup>1</sup> Induced growth inhibition, G0/G1 arrest and apoptosis in a variety of cancer cell lines, with a B-Raf mutation favoring but not guaranteeing a response.<sup>2</sup> Inhibits the growth of B-Raf V600E-positive melanomas *in vitro* and *in vivo*.<sup>3</sup> Treatment of patients that carry the V600E BRAF mutation resulted in complete or partial tumor regression.<sup>4</sup>

- 1) Khazak et al. (2007), Selective Raf inhibition in cancer therapy; Expert Opin. Ther. Targets, 11 1587
- 2) Tap et al. (2010), Pharmacodynamic characterization of the efficacy signals due to selective BRAF inhibition with PLX4032 in malignant melanoma; Neoplasia, **12** 637
- 3) Lee et al. (2010), PLX4032, a potent inhibitor of the B-Raf V600E oncogene, selectively inhibits V600E-positive melanomas; Pigment Cell Melanoma Res., **23** 820
- 4) Flaherty et al. (2010), Inhibition of mutated, activated BRAF in metastatic melanoma; N. Engl. J. Med., 363 809

## PHYSICAL DATA

Molecular Weight:	489.92
Molecular Formula:	C <sub>23</sub> H <sub>18</sub> CIF <sub>2</sub> N <sub>3</sub> O <sub>3</sub> S
Purity:	99% by HPLC
Solubility:	DMSO (100mg/mL)
Physical Description:	White to Off-white solid
Storage and Stability:	Store as supplied desiccated at -20°C for up to 1 year from the date of purchase. Solutions in
	DMSO may be stored at -20°C for up to 3 months.

Materials provided by Focus Biomolecules are for laboratory research use only and are not intended for human or veterinary applications.

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