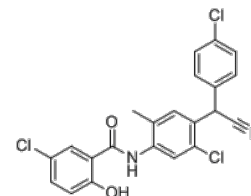


Product Name : ZT-1a
Cat. No. : PC-72593
CAS No. : 212135-62-1
Molecular Formula : C₂₂H₁₅Cl₃N₂O₂
Molecular Weight : 445.73
Target : Other Targets
Solubility :



Biological Activity

ZT-1a (STK39 inhibitor ZT-1a, SPAK-IN-ZT-1a) is a novel potent, selective SPAK kinase (STK39) inhibitor.

ZT-1a potently inhibited SLC12A cation-Cl-cotransporters (CCC), phosphorylation of NKCC1 p-Thr203/207/212 by 72% at 1uM and phosphorylation of KCC sites 1/2 by 65-77% at 3 uM in HEK-293 cells.

ZT-1a is a more potent modulator of SPAK-dependent CCC phosphorylation than the current SPAK kinase inhibitors Closantel, STOCK1S-50699, and STOCK1S-14279.

ZT-1a inhibited SPAK phosphorylation at Ser373 in HEK-293 cells (70% inhibition at 3-10 uM).

ZT-1a disrupts SPAK interaction with WNK but not with MO25 α , reduces SPAK-dependent CCC phosphorylation in cells, inhibits NKCC1 but stimulates KCC3 activity.

ZT-1a decreases SPAK-CCC phosphorylation in hemorrhagic hydrocephalus, modulates primary cortical neuron volume, reduces stroke-associated cerebral edema and infarct, improves neurological function after ischemic stroke in vivo.

References

Jinwei Zhang, et al. Nat Commun. 2020 Jan 7;11(1):78.

Caution: Product has not been fully validated for medical applications. Lab Use Only!

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