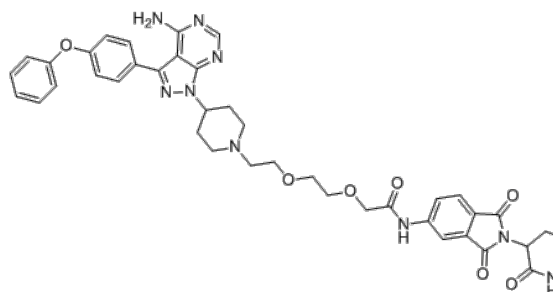


**Product Name** : MT-802  
**Cat. No.** : PC-35213  
**CAS No.** : 2231744-29-7  
**Molecular Formula** : C<sub>41</sub>H<sub>41</sub>N<sub>9</sub>O<sub>8</sub>  
**Molecular Weight** : 787.834  
**Target** : PROTAC  
**Solubility** :



## Biological Activity

MT-802 (MT802) is a potent **BTK PROTAC** that induces degradation of both wild-type and C481S mutant BTK (DC50=9.1 nM); MT-802 recruits BTK to the cereblon E3 ubiquitin ligase complex to trigger BTK ubiquitination and degradation via the proteasome; MT-802 binds fewer off-target kinases than ibrutinib does and retains an equivalent potency (>99% degradation at nanomolar concentrations) against wild-type and C481S BTK, elicits complete BTK knockdown at 250 nM; reduces the pool of active, phosphorylated BTK in cells isolated from CLL patients with the C481S mutation, whereas ibrutinib cannot.

## References

Buhimschi AD, et al. *Biochemistry*. 2018 Jun 14. doi: 10.1021/acs.biochem.8b00391.

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

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