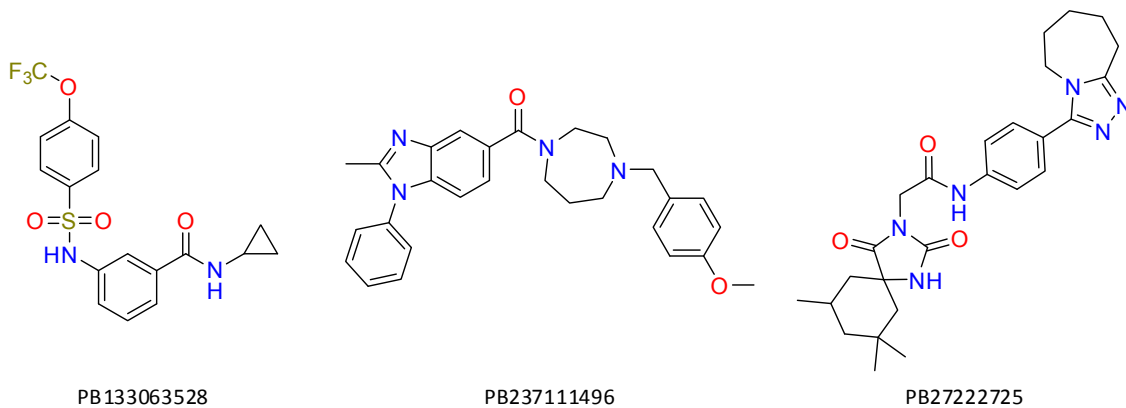


UORSY PPI Modulators

Modulating protein-protein interactions (PPIs) has been considered as an important goal for drug discovery because of their role in numerous cell disorders.¹ PPIs, however, are both highly promising and highly difficult to study.² Much of the success in finding “hidden gems” – small molecule modulators of PPIs, – relies on the quality of the starting library. Understanding this, we created a set of PPIs modulators utilizing the following approaches:

- 1) physicochemical criteria (“Rule of 4” for inhibitors of PPIs);
- 2) the “hot spot” paradigm (design included functional groups able to interact with Tyr, Trp, Arg residues);³
- 3) ligand-based design (similarity analysis to TIMBAL database).⁴



Physicochemical profiles of **UORSY PPI modulators**:

$400 < \text{MW} < 600$; $2 < \text{HbA} < 10$; $0 < \text{HbD} < 4$; $2 < \log P < 6$; $1 \leq \text{RotBonds} \leq 12$; $\text{TPSA} > 50$.

UORSY PPI modulators are available in stock and could be delivered within 2 weeks in any customer-preferred format: as powders, dry films or DMSO solutions formatted in vials, 96 or 384-well plates. All compounds have a minimum purity of 90% assessed by ^1H NMR; analytical data is provided.

For more information, please contact us at screenlibs@uorsy.com

¹Morelli, X.; Hupp, T., *EMBO Rep.* **2012**, 13, 877–879

²Surade, S.; Blundell, T. L., *Chemistry and Biology*, **2012**, 19, 42–50

³Wells, J. A.; McClendon, C. L., *Nature* **2007**, 450, 1001–1009

⁴Higuero, A. P.; Jubbe, H.; Blundell, T. L., *Database (Oxford)*. **2013**, 2013, bat039