

PHARMACOLOGICAL CHARACTERIZATION OF NEW ANTAGONIST OF THE EPH RECEPTORS ENDOWED WITH ANTI-ANGIOGENIC ACTIVITY

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Eph receptors and ephrin ligands are overexpressed in a large number of cancer types, such as ovarian and breast cancer [1-2] where they promote metastasis, invasion and tumor angiogenesis [3]. For this reason, targeting the Eph-ephrin system could be a new approach in cancer therapy.

The activity of the Eph receptors can be inhibited by means of intracellular kinase inhibitors or by using protein-protein inhibitors, targeting the extracellular ligand-binding domain of Eph receptor which include antibodies, chimeric proteins and small molecules [4].

Our research group focused its attention on the development of protein-protein inhibitors and we recently obtained UniPR129 [5], (the conjugate of lithocholic acid with L-homo-Tryptophan). It was a competitive and reversible antagonist of EphA2 receptor, endowed with high potency in in vitro anti-angiogenic assays but endowed with unfavourable physicochemical properties which limited its use in vivo.

To overcome this issue, metadynamics was employed to design new derivatives [6] and UniPR139 and UniPR502 emerged as a couple of the most promising compounds.

They showed a K_i of 950 nM and 750 nM respectively blocking the Eph-ephrin interaction in a competitive and reversible manner. They did not discriminate Eph receptor subclasses and they reduced ephrin-A1-induced EphA2 receptor phosphorylation in the low micromolar range without exerting non specific cytotoxic effects. The compounds showed an in vitro and in vivo anti-angiogenic activity, when tested in tube formation assay on HUVEC and in CAM assay, at 3-10 μ M concentrations.

The anti-angiogenic activity was related to a reduction of cell proliferation and VEGF-induced VEGFR2 activation, in fact at 30 μ M UniPR139 and UniPR502 partially interfere with the VEGFR phosphorylation and internalization

IN conclusion UniPR139 and UniPR502 are promising pharmacological for further in vivo characterization.

References:

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