

## Nové HSD17beta1 inhibitory pro léčbu rakoviny

### Fáze vývoje technologie

#### Preklinické testy

### Status IP ochrany

PCT/CZ2017/050022

### Strategie pro hledání partnera

Licencování, Spolupráce

### Instituce

Ústav organické chemie a biochemie AV ČR, v.v.i.

### Vlastník

IOCB, IMG Pague, IMTM  
Olomouc and Helmholtz Zenter  
Munich

### Motivace

17 $\beta$ HSD1 is an enzyme that has been known to play a pivotal role in tissue specific estradiol (E2) synthesis for more than fifty years. E2 is a very potent hormone, which regulates the expression of a variety of genes by binding to estrogen receptors (ER) and thus it plays a crucial role in the physiological as well as pathological proliferation and differentiation of the target cell. 17 $\beta$ HSD1 affects breast cancer cell proteome and modulates expression of several genes at both mRNA and protein levels. Furthermore, it is associated with an increased risk of cell migration and cancer relapse. Therefore regulation of 17 $\beta$ HSD1 should be considered as potential novel endocrine therapy or 17 $\beta$ HSD1 expression as an independent prognostic marker in breast cancer patients.

### Popis

We have successfully developed highly potent and specific non-estrogenic inhibitors of 17 $\beta$ HSDs. We have recently shown the efficacy of these compounds in in vitro and in vivo testing of E1 to E2 conversion after application of lead compounds. SAFETY - NOVELTY No toxic effect up to 2mg/kg Low estrogenicity of lead compound EP449 Experiments on T47D cells show inhibition of HSD17b1 (accumulation of E1 in HPLC MS/MS detectable) proof of concept experiment in eggs has been successful - with reduced tumor size and number of metastases, in comparison to Tamoxifen (standard care)

### Komerční využití

We are looking for out-licensing partners or partners to co-develop the project further