

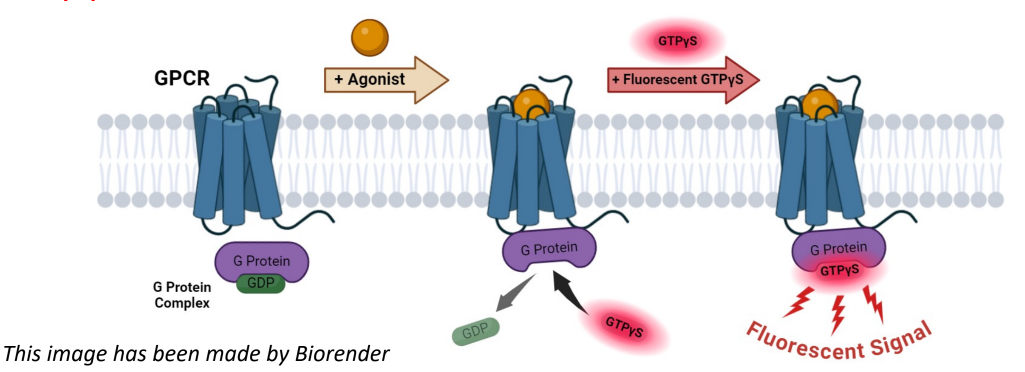
Innovative Chemistry to Innovative Chemistry to Innovative Biology Fluorescent GTPyS

GPCR Functional assay

GPCR functional activity data are extremally important in Drug Discovery programs, both for monitoring therapeutical effects or off target side effects.

A reliable and efficient assay is based on the use of no-hydrolysable GTP derivative, GTP \(\text{S} \), which allows to detect the activation of G protein after ligand binding.

Compared to the radioactive [35]GTPγS, fluorescent GTPγS enables assays to be performed with the same sensitivity and robustness while allowing the use of conventional facilities and equipment



Fluorescent GTP_VS

In the frame of a collaboration between Celtarys and Arcoscreen, we developed a **Fluorescent GTPyS** (CELT-503) emitting the in red region of the spectra (589/616).

The developed fluorescent probe was validated by Arcoscreen in its own platform for GPCRs functional screening.







Celtarys

Advantages of Fluorescent GTPγS vs conventional radioactive GTPγS

	[³⁵S]GTPγS	Fluorescent GTPγS
No-hydrolysable GTP derivative	√v	√v
Detect the direct activation of G protein, not second messengers	√v	√v
Special handling	YES (Radioactive)	NO
Compatible with common equipment	×x	√v
Compatible with High Throughput Screening (HTS)	×x	√v
Costs per assay	↑	\downarrow

As a radioactive substance, [35]GTPyS requires additional precautions, including trained personnel and specific safety measures, as well as specialized facilities for handling radioactive materials like sulfur-35. Additionally, the radioactive waste generated during the assays must be managed and disposed of appropriately. Furthermore, the [35S] isotope has a relatively short half-life of 87 days, compared to the 12-year half-life of tritium.

These challenges are effectively addressed by replacing radioactivity with fluorescence (CELT-503), offering a safer and more practical alternative.

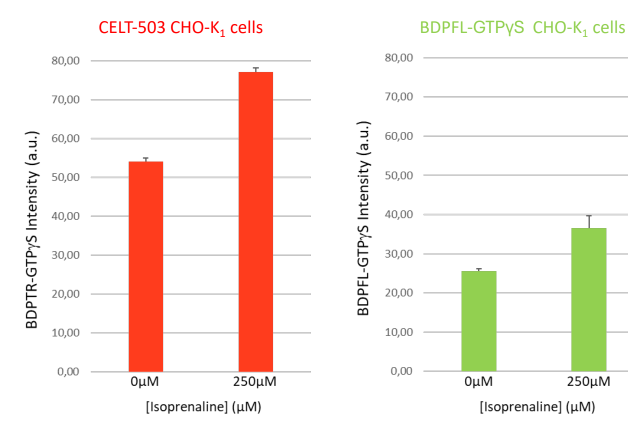
GPCRs Functional assay

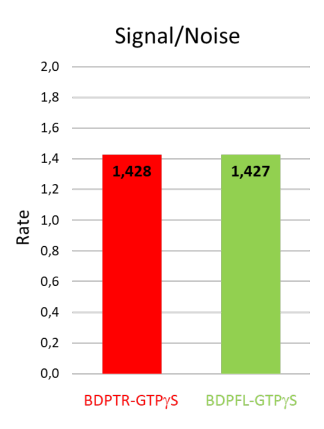
α-adrenergic receptors

Experiment performed in triplicate in CHO-K1 cells (passage 12) expressing endogenously adrenoreceptors. Experiment read-out is the zeiss fluorescent microscope axiovert 7 with the following parameters:

- -CELT-503: ex. 555 nm, em 614 nm, 50% intensity light source, 150ms exposure time
- -BDPFL-GTPγS: ex. 475 nm, em 509 nm , 50% intensity light source, 500ms exposure time

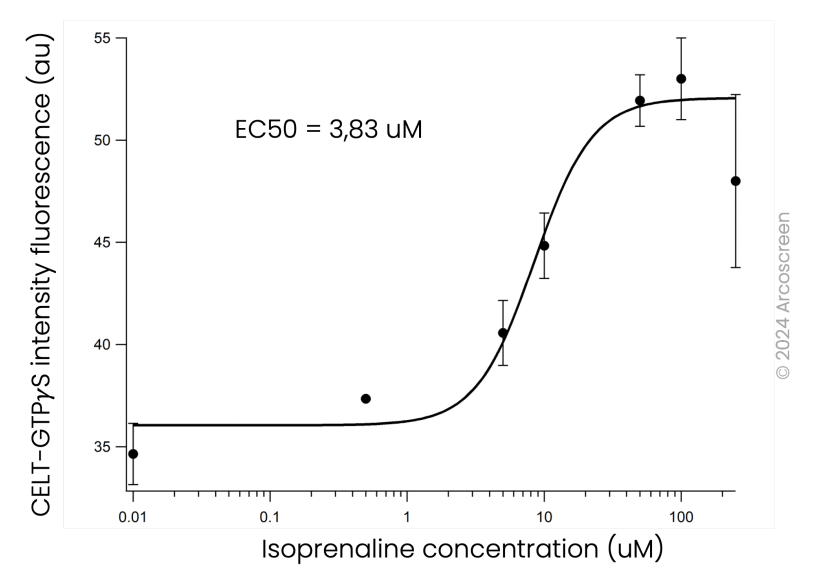
Compared to the BDPFL-GTPγS (actual market available), the CELT-503 (Celtarys) is 2x brighter with 3 times less exposure time, while keeping the same ratio Signal/Noise. As it emits in the red spectrum, it is compatible with most multiplexing experiments, as most fluorophores emit in the green spectrum.





GPCR Dose-response curve

The EC_{50} of isoprenaline was measured using CELT-503. The experiments were performed in triplicates in CHO-K1 cells endogenously expressing adrenoreceptors.





Thank you

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