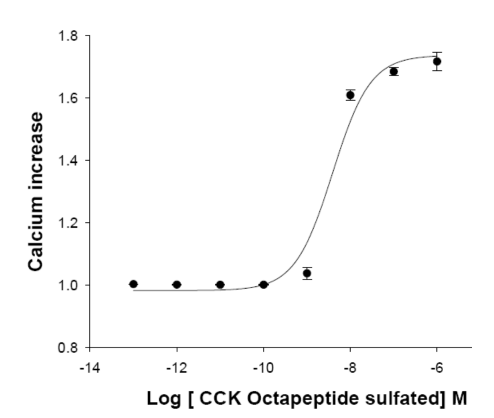


HiTSeeker CELL LINES (LABEL-FREE GPCRS)

- CHOLECYSTOKININ B RECEPTOR (CCK2) CELL LINE -



Product name: CCK₂ (CCKBR) /U2OS cell line Ec₅₀ CCK Octapeptide, sulfated: 3.9 x 10⁻⁹ M

Z′: 0.87+/- 0.02



REF: P30179

- CHOLECYSTOKININ B RECEPTOR (CCK2) U2OS CELL LINE -

Product Name: CCK₂ (CCKBR)/U2OS

Official Full Name: Cholecystokinin A receptor

DNA Accession Number: GenBank: AY322551

Host Cell: U2OS

Format: 2 cryopreserved vials

Resistance: G418

Size: $> 3 \times 10^6$ cells / vial

Storage: Liquid Nitrogen

📀 Assay Briefly description

Each vial of HiTSeeker CCK₂/U2OS contains U2OS cells stably expressing human Cholecystokinin B receptor (CCK₂) with no tag.

Innoprot HiTSeeker CCK_2 cell line has been designed to assay compounds or analyze their capability to modulate Cholecystokinin B receptor. When the agonist binds to CCK_2 a G protein is activated, which in turn, triggers a cellular response mediated by second messengers (Calcium).

This cell line has been validated measuring calcium increase in the cytosol. The high reproducibility of this assay allows monitoring CCK_2 activation process in High Throughput Screening.

About CCK₂

CCK receptors family is composed of two GPCRs known as CCK₁ and CCK₂ receptors. Both receptors bind Cholecystokinin (CCK) that is a main gastrointestinal and neuronal peptide hormone, involved in stimulating gallbladder contraction, pancreatic secretion, gastrointestinal motility and satiety.

The CCK₁₂ receptor has a high expression in several tumour types including medullary thyroid carcinoma (MTC), neuroendocrine tumours, small cell lung cancer, and colorectal cancers.



🔊 Assay Characterization

Our expression plasmid contains the coding sequence of human CCK₂ protein. Our plasmid was transfected in U2OS cells. Resistant clones were obtained by limit dilution and receptor gene expression was tested by RT-PCR using GAPDH as internal control (Fig.1).

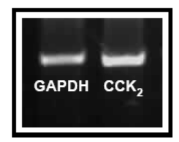


Fig.1. GAPDH housekeeping gene and CCK2, RT-PCR.

S Validation of CCK₂ cell line

Calcium assay (Ec50 = 3.9 x 10⁻⁹M)

A typical fluorescent calcium assay was performed using Fura-2/AM ratiometric. Calcium increase inside the cell was measured using the ratio of the fluorescence from Fura2 bound and not bound to the ion. Image acquisition was performed using a "BD Pathway 855" High-Content Bioimager from BD Biosciences.

Cells were incubated with Fura2-AM and treated with increasing CCK Octapeptide, sulfated concentrations.

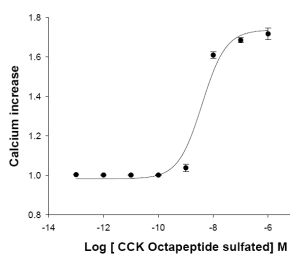


Fig.2. CCK_2 dose response in calcium assay. Cells were treated with CCK Octapeptide, sulfated concentrations ranging from 0 to 1 μ M, n=5. The EC50 for **CCK Octapeptide**, sulfated was ~ **3.9×10⁻⁹M**. The calcium assay was validated with a Z'=0.87+/-0.02 for High Content Screening.