

Research use only. Not for use in diagnostic procedures.

ValiScreen® GPCR Cell Line

# human Somatostatin Receptor sst<sub>4</sub> Cell Line

Product No.: ES-524-C Lot No.: 1893891

Material Provided

Cells: 2 x 1 mL frozen aliquot (ES-524-CV)

Format: ~2.5 x 10° cells /mL in freezing medium

**Product Information** 

Cellular Background: CHO-K1

Cell Line Development: Our proprietary bicistronic expression plasmid containing the

sequence coding for the human Somatostatin receptor sst<sub>4</sub> was transfected in CHO-K1 cells. Geneticin-resistant clones were obtained by limit dilution and compared for receptor expression levels using a radioligand binding assay. The clone with the highest receptor expression level was selected for characterization in

binding and functional assays.

DNA Sequence: Identical to coding sequence of GenBank L14856.1.

Corresponding Protein Sequence: Identical to GenBank AAA36623.1.

Receptor expression level ( $B_{MAX}$ ): Estimated to be 13.8  $\pm$  7.0 pmol/mg protein, using [ $^{125}I$ ]-[Tyr $^{11}$ ]-SRIF-

14

 $K_D$  for the above radioligand: 1.9  $\pm$  0.14 nM

Shipping Conditions: Shipped on dry ice. Please ensure dry ice is still present in the

package upon receipt or contact customer support.

Storage Conditions: Store in liquid nitrogen (vapor phase) immediately upon receipt.



# **Quality Control**

The EC $_{50}$  for a reference agonist was determined in LANCE $^{\circ}$  Ultra cAMP assay performed on an EnVision $^{\circ}$  instrument. A mycoplasma test was performed using MycoAlert $^{\circ}$  (Lonza) mycoplasma detection kit. We certify that these results meet our quality release criteria.

SRIF-28 (EC<sub>50</sub>): 0.26 nM

Stability: Cells were kept in continuous culture for at least 60 days and showed no

decrease of receptor expression level in a saturation binding assay (stable  $B_{max}$  and  $K_d$ ) and no decrease in functional response (EC<sub>50</sub>,  $E_{max}$  in cAMP

assay).

Mycoplasma: This cell line tested negative for mycoplasma.

# **Assay Procedures**

We have shown for many of our GPCR cell lines that freshly thawed cells respond with the same pharmacology as cultured cells. All of our products validated in this way are available as frozen ready-to-use cells in our catalogue. This demonstrates that cells can be prepared and frozen in advance of a screening campaign simplifying assay logistics.



## Recommended Cell Culture Conditions (CHO-K1)

- The recommended media catalogue number and supplier reference information are listed in this Product Technical Data Sheet (last page). Media composition is specifically defined for each cell type and receptor expression selection. The use of incorrect media or component substitutions can lead to reduced cell viability, growth issues and/or altered receptor expression.
- Cells undergo major stress upon thawing, and need to adapt to their new environment which may initially affect
  cell adherence and growth rates. The initial recovery of the cells, and initial doubling time, will vary from
  laboratory to laboratory, reflecting differences in the origin of culture media and serum, and differences in
  methodology used within each laboratory.
- For the initial period of cell growth (i.e. until cells have reached Log-phase, typically 4-10 days), we strongly recommend removal of the antibiotics (G418, Zeocin™, Puromycin, Blasticidin, Hygromycin, Penicillin and Streptomycin) from the culture media. Immediately after thawing, cells may be more permeable to antibiotics, and a higher intracellular antibiotic concentration may result as a consequence. Antibiotics should be reintroduced when cells have recovered from the thawing stress.

Growth Medium: Ham's F-12, 10% FBS, 0.4 mg/ml G418 (receptor expression selection).

Freezing Medium: Ham's F-12, 10% FBS with 10% DMSO, without selection agents.

Thawing Cells: Using appropriate personal protective equipment, rapidly place the frozen aliquot in a  $37^{\circ}$ C water bath (do not submerge) and agitate until its content is thawed completely. Immediately remove from water bath, spray aliquot with 70% ethanol and wipe excess. Under aseptic conditions using a sterile pipette, transfer content to a sterile centrifuge tube containing 10 mL growth medium without antibiotics, pre-warmed at  $37^{\circ}$ C, and centrifuge ( $150 \times g$ ,  $5 \times g$ ,  $5 \times g$ ). Discard supernatant using a sterile pipette. Resuspend cell pellet in  $10 \times g$  for  $10 \times g$ ,  $10 \times$ 

Recommended Seeding Density: Thawing: 15000 - 33000 cells/cm<sup>2</sup>

Log-phase: 11000 - 15000 cells/cm<sup>2</sup>

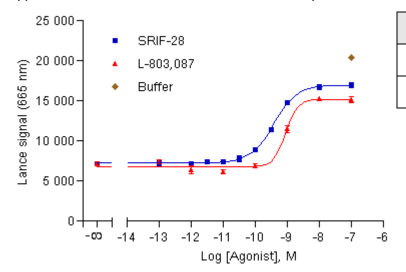
Troubleshooting: Initial doubling time can vary between 18 and 96 hours (Average = 25 hours). If cells are still not adhering after 48 hours or grow very slowly, we recommend maintaining the cells in culture and not replacing the media before 5-6 days (cells secrete factors that can help with adherence and growth). If confluence is still <50% after 5-6 days, it is recommended that you replace the media with fresh media (without antibiotics). Do not passage the cells until they reach 80-90% confluence (Log-phase). If cells have not recovered after 10-12 days, please contact our Technical Support.

Culture Protocol: Under aseptic conditions, cells are grown to 80% confluence (Log-phase) and trypsinized (0.05% trypsin/0.5 mM EDTA in calcium and magnesium-free PBS). See recommended seeding density for Log-phase above.

Banking Protocol: Cells are grown to 70-80% confluence (Log-phase). Under aseptic conditions, remove medium and rinse the flask with an appropriate volume of calcium and magnesium-free PBS (example 10 mL for T-175). Trypsinize (0.05% trypsin/0.5 mM EDTA in calcium and magnesium-free PBS) to detach cells (example 5 mL for T-175), let stand 5-10 min at 37°C. Add fresh, room temperature growth medium (without antibiotics) to stop trypsinization and dilute EDTA (example 10 mL for T-175). Transfer cells to a sterile centrifuge tube and centrifuge (150 x g, 5 min). Discard supernatant using a sterile pipette. Resuspend cell pellet in ice-cold freezing medium by pipetting up and down to break up any clumps. Count cells and rapidly aliquot at the selected cell density (e.g. 2.5 x  $10^6$  cells/mL) in sterile polypropylene cryovials. Use appropriate material to ensure slow cooling (about -1°C/min) until -70°C. Transfer vials into a liquid nitrogen tank (vapor phase) for storage.



# Typical Product Data -LANCE® cAMP Assay



| Agonist   | EC <sub>50</sub> (M)    |  |
|-----------|-------------------------|--|
| SRIF-28   | 3.9 x 10 <sup>-10</sup> |  |
| L-803,087 | 9.0 x 10 <sup>-10</sup> |  |

Figure 1. Agonist Response in LANCE® cAMP assay

An agonist dose-response experiment was performed in 384-well format using 2 500 cells/well. Cells were thawed and incubated for 30-min with 10  $\mu$ M Forskolin (FK) and the indicated agonist concentrations. Cells incubated in the absence of agonist and FK were included as a control (Buffer). Time-resolved fluorescence was measured on an EnVision® instrument. Data from a representative experiment are shown.



Typical Product Data -Radioligand Binding Assay (Filtration)

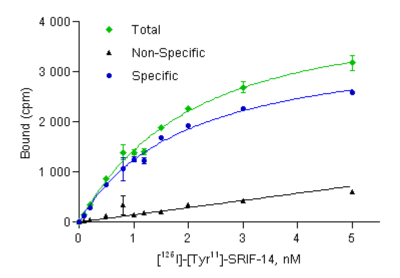
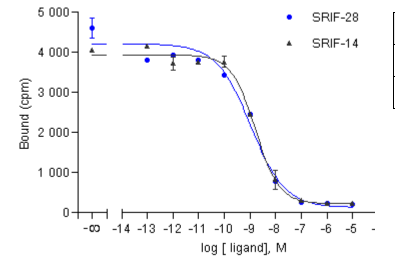


Figure 2: Saturation Binding Assay Curve (Filtration)
A saturation binding assay was performed in 96-well format using 0.8 µg membranes/well. Counts per minute (cpm) were measured on a TopCount® instrument. Data from a representative experiment are shown.

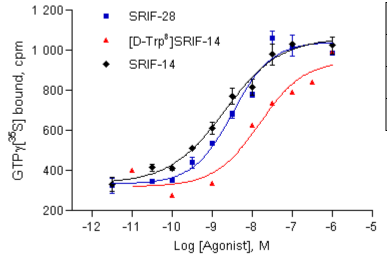


| Agonist / Antagonist | IC <sub>50</sub> (M)   |  |
|----------------------|------------------------|--|
| SRIF-28              | 1.1 x 10 <sup>-9</sup> |  |
| SRIF-14              | 1.6 x 10 <sup>-9</sup> |  |

Figure 3: Competition Binding Assay Curve (Filtration)
A competition binding assay was performed in 96-well format using 0.8 µg membranes/well. Displacement of 0.2 nM [1251]-[Tyr11]-SRIF-14 was used. Counts per minute (cpm) were measured on a TopCount® instrument. Data from a representative experiment are shown.



Typical Product Data - GTPγS - SPA® Assay



| Agonist                      | EC <sub>50</sub> (M)   |
|------------------------------|------------------------|
| SRIF-28                      | 3.2 x 10 <sup>-9</sup> |
| SRIF-14                      | 2.1 x 10 <sup>-9</sup> |
| [D-Trp <sup>8</sup> ]SRIF-14 | 1.5 x 10 <sup>-8</sup> |

Figure 4. Agonist Response in GTP $\gamma$ S - SPA® assay An agonist dose-response scintillation proximity assay (SPA) was performed in 96-well format using 5  $\mu$ g membranes/well. Counts per minute (cpm) were measured on a TopCount® instrument. Data from a representative experiment are shown.



## LANCE® Ultra cAMP Assay Procedure

Stimulation Buffer: HBSS, 5 mM HEPES, 0.1 % Protease-free BSA, 0.5 mM IBMX, pH 7.4.

Cells/well: For compounds not tested herein we recommend titrating the cells for optimal

performance, i.e. 500-3000 cells per assay point.

cAMP measurements can be performed with the LANCE® *Ultra* cAMP 384 Kit (Revvity # TRF0262), according to the manufacturer instructions. Briefly:

Protocols for a 384-well white Optiplate (total assay volume of 20  $\mu$ L):

| cAMP<br>Standard curve  | G <sub>s</sub> Agonist  | G <sub>s</sub> Antagonist | G <sub>i</sub> Forskolin<br>titration | G <sub>i</sub> Agonist  | G <sub>i</sub> Antagonist   |
|---|-------------------------|---------------------------|---------------------------------------|-------------------------|-----------------------------|
| 5 µL cAMP<br>Standard   | 5 µL cell<br>suspension | 5 µL cell<br>suspension   | 5 µL cell<br>suspension               | 5 µL cell<br>suspension | 5 µL cell<br>suspension     |
| 5 μL<br>Stimulation<br>Buffer   | 5 μL Agonist            | 2.5 µL<br>Antagonist      | 5 μL Forskolin                        | 2.5 µL<br>Agonist       | 2.5 µL Antagonist           |
| -   | -                       | 2.5 µL Agonist            | -                                     | 2.5 µL<br>Forskolin     | 2.5 µL<br>Forskolin/Agonist |
| Incubate 30 min at room temperature (optional step for cAMP Standard curve) |                         |                           |                                       |                         |                             |
| 5 μL 4X Eu-cAMP Tracer Working Solution                                     |                         |                           |                                       |                         |                             |
| 5 μL 4X U <i>Light</i> -anti-cAMP Working Solution                          |                         |                           |                                       |                         |                             |
| Incubate 1 h at room temperature  |                         |                           |                                       |                         |                             |
| Read on an EnVision® instrument. Remove microplate seal prior to reading    |                         |                           |                                       |                         |                             |

- 1. Cells in mid-log phase, grown in media without antibiotics for 18 hours prior to the experiment, are detached by gentle flushing with PBS-EDTA, recovered by centrifugation and resuspended in stimulation buffer e.g. at the concentration of  $6.0 \times 10^5$  cells/mL (for 3000 cells/well).
- 2. Prepare the 4X Tracer Working Solution by making a 1/50 dilution of the Eu-cAMP stock solution in the cAMP Detection Buffer.
- 3. Prepare an ULight-anti-cAMP Intermediate Solution by making a 1/10 dilution of the ULight-anti-cAMP stock solution in cAMP Detection Buffer. Prepare the 4X ULight-anti-cAMP Working Solution by making a 1/30 dilution of the ULight-anti-cAMP intermediate solution in the cAMP Detection Buffer.

#### Notes:

For 96- and 1536-well formats, adjust proportionally the volume of each assay component in order to maintain the volume ratios for the 384-well format. Do not modify the Eu-cAMP and/or the ULight-anti-cAMP concentrations.



### LANCE® cAMP Assay Procedure

Stimulation Buffer: HBSS, 5 mM HEPES, 0.1 % Protease-free BSA, 0.5 mM IBMX, pH 7.4.

Cells/well: 2500. For compounds not tested herein we recommend titrating the cells for

optimal performance, i.e. 1000-10000 cells per assay point.

Antagonist Pre-incubation: Simultaneous addition of antagonists with reference agonist.

Agonist Stimulation: 30 min at room temperature (22°C).

cAMP measurements were performed with the LANCE® cAMP 384 Kit (Revvity # AD0262), according to the manufacturer instructions. Briefly:

1. Compounds (6  $\mu$ L/well) were dispensed into a 384-well white Optiplate:

|            | $G_{\alpha s}$ and $G_{\alpha i}$ assay modes |                        | G <sub>αs</sub> assay mode |                        | G <sub>αi</sub> assay mode                  |   |
|------------|---|------------------------|----------------------------|------------------------|---|---|
|            | Basal   | Forskolin              | Agonist<br>Assay           | Antagonist<br>Assay    | Agonist Assay                               | Antagonist Assay                                  |
| Buffer     | 6 μL  | -                      | -                          | -                      | -   | -   |
| Antagonist | -   | -                      | -                          | 3 μL of 4x final conc. | -   | 3 μL of 4x final conc.                            |
| Agonist    | -   | -                      | 6 μL of 2x<br>final conc.  | 3 μL of 4x final conc. | 6 μL of 2x final conc. in 2x final FK conc. | 3 μL of 4x final<br>conc. in 4x final<br>FK conc. |
| Forskolin  | -   | 6 μL of 2x final conc. | -                          | -                      | FR CONC.                                    | FR conc.  |

- 2. Cells in mid-log phase, grown in media without antibiotics for 18 hours prior to the experiment, were detached by gentle flushing with PBS-EDTA, recovered by centrifugation and resuspended in stimulation buffer at the concentration of  $4.2 \times 10^5$  cells/mL.
- 3. The Alexa Fluor® 647-anti cAMP antibody was added 1/100 (vol/vol) to the cell suspension.
- 4.  $6 \mu L/well$  of cell and antibody suspension (2500 cells/well) were dispensed on top of the compounds prepared in the 384 well Optiplate.
- 5. After incubation for 30 min at room temperature the reaction was stopped by addition of 12  $\mu$ L of Detection Mix.
- 6. The plate was incubated for 60 min at room temperature and read on an EnVision®.

Note: Assays can also be miniaturized into 1536-well format.



# Membrane Radioligand Binding Assay Procedure (Filtration)

Note: The following are recommended assay conditions and may differ from the conditions used to generate the typical data shown in the above section.

Assay Buffer: 25 mM HEPES pH 7.4, 10 mM MgCl<sub>2</sub>, 1 mM CaCl<sub>2</sub>, 0.5% BSA

Wash Buffer: 50 mM Tris-HCl pH 7.4 at 4°C, 0.2% BSA

Radioligand: [125I]-[Tyr11]-SRIF-14 (Revvity # NEX446)

Filters: Unifilter 96 GF/C (Revvity # 6055690)

Membrane Binding Protocol:

Binding assays were performed in 200  $\mu L$  total volume according to the following conditions. All dilutions are performed in assay buffer:

| 1.                   | Membrane dilution:                                     | 0.25 μg of membranes per well, diluted in order to dispense 150μL/well. Keep on ice.  |  |  |
|----------------------|--|---|--|--|
| ,                    | Assembly on ice n 96 Deep well plate) uration Binding: | <ul> <li>25 μL of assay buffer or of unlabeled ligand (Somatostatin-28, 1 μM final) for determination of non specific binding</li> <li>25 μL of radioligand at increasing concentrations (see figure 2)</li> <li>150 μL of diluted membranes</li> </ul> |  |  |
| Competition Binding: |  | <ul> <li>25 μL competitor ligand at increasing concentrations (see figure 3)</li> <li>25 μL of radioligand (0.4 nM final)</li> <li>150 μL of diluted membranes</li> </ul>   |  |  |
| 3.                   | Incubation:  | 60 min at 27°C.   |  |  |
| 4.                   | Filters preparation:                                   | GF/C filters were presoaked in 0.5 % PEI at room temperature for at least 30 min.   |  |  |
| 5.                   | Filtration:  | Aspirate and wash 9 x 500 µL with ice cold wash buffer using a FilterMate Harvester.  |  |  |
| 6.                   | Counting:  | Add 30 µL/well of MicroScint™-O (Revvity # 6013611), cover filter with a TopSeal-A PLUS (Revvity # 6050185) and read on a TopCount®.  |  |  |



# GTPγS - SPA® Assay Procedure

Assay Buffer: 20 mM HEPES pH 7.4, 100 mM NaCl, 10  $\mu$ g/ml saponin, 10 mM MgCl<sub>2</sub>, 0.1% protease-

free BSA

GDP concentration: 10 µM GDP (final)

SPA Beads: PVT-WGA (Revvity # RPNQ0001), 0.25 mg/well

Radioligand: GTP $\gamma$ S, [ $^{35}$ S] - (Revvity # NEG030H)

Membranes: 5 μg/well

Format: 96-well

Final volume:  $100 \mu L/well$ 

 $GTP\gamma S\text{-SPA assays were performed in 100 }\mu L \text{ total volume according to the following conditions. All dilutions are}$ 

performed in assay buffer:

| <u>seriorme</u> | ed in assay buffer:                      |  |  |  |
|-----------------|--|--|--|--|
| 1.              | Membrane Dilution:                       | 5 µg of membranes per well, diluted in order to dispense 10 µL/well. Keep on ice.  |  |  |
| 2.              | GDP saturation:                          | Mix a 10-fold concentrated GDP solution (i.e. 100 μM) with the membranes dilution. Incubate "membranes : GDP mix" on ice for 15 min.   |  |  |
| 3.              | GTPγS, [ <sup>35</sup> S] - dilution:    | Dilute GTPγS, [ <sup>35</sup> S] - to give ~25.000 dpm/10μL  |  |  |
| 4.              | Beads:                                   | Dilute beads to 25 mg/mL (0.25 mg/10 $\mu$ L). Premix beads with the GTP $\Box$ S, [ $^{35}$ S] - dilution just before starting the reaction ("GTP $\gamma$ S, [ $^{35}$ S] - : Beads mix").   |  |  |
|                 | Assembly (in Optiplate™),<br>nist Assay: | <ul> <li>50 μL of 2x agonist dilution at increasing concentrations</li> <li>20 μL of the "membranes : GDP mix"</li> <li>10 μL of assay buffer</li> <li>20 μL of the "GTP γS, [35S] - : Beads mix"</li> </ul>   |  |  |
| Anta            | agonist Assay:                           | <ul> <li>50 μL of 2x antagonist dilution at increasing concentrations</li> <li>20 μL of the "membranes: GDP mix"</li> <li>10 μL of 10x reference agonist dilution to reach a final concentration corresponding to its EC<sub>80</sub></li> <li>20 μL of the "GTPγS, [35S] -: Beads mix"</li> </ul> |  |  |
| 6.              | Incubation:                              | <ul> <li>Cover plate with a TopSeal</li> <li>Shake on an orbital shaker for 2 min</li> <li>Incubate for 1h at RT°</li> <li>Centrifuge the plate for 10 min at 2000 rpm</li> <li>Incubate for 0h to 4h at RT°</li> </ul>  |  |  |
| 7.              | Counting                                 | Count for 1 min on a TopCount®   |  |  |



### References

- 1. Demchyshyn LL, Srikant CB, Sunahara RK, Kent G, Seeman P, Van Tol HH, Panetta R, Patel YC, Niznik HB. (1993) Cloning and expression of a human somatostatin-14-selective receptor variant (somatostatin receptor 4) located on chromosome 20. Mol Pharmacol. 43:894-901.
- 2. Xu Y, Song J, Bruno JF, Berelowitz M. (1993) Molecular cloning and sequencing of a human somatostatin receptor, hSSTR4. BBRC. 193:648-652.
- 3. Siehler S, Hoyer D (1999) Characterisation of human recombinant somatostatin receptors. 4. Modulation of phospholipase C activity. Naunyn Schmiedebergs Arch Pharmacol. 360:522-532.
- 4. Csaba Z, Dournaud P (2001) Cellular biology of somatostatin receptors. Neuropeptides. 35:1-23.



#### Materials and Instrumentation

The following tables provide the references of compounds and reagents used or recommended for the characterization of the human Somatostatin receptor sst<sub>4</sub> ValiScreen® cell line, as well as some advice on how to use these compounds:

Table 1. References of compounds used for functional characterization and binding assays

| Name                           | Provider | Cat no | Working Stock Solution                 |
|--------------------------------|----------|--------|--|
| SRIF-28 (Somatostatin-28)      | Bachem   | H-4955 | 200 μM in PBS / 0.1% protease-free BSA |
| SRIF-14 (Somatostatin-14)      | Bachem   | H-1490 | 200 μM in 1% Acetic Acid               |
| [D-Trp <sup>8</sup> ]SRIF-14   | Bachem   | H-3198 | 1 mM in PBS / 0.1% protease-free BSA   |
| L-803,087                      | Tocris   | 1979   | 10 mM in DMSO                          |
| [125][Tyr11]-Somatostatin-14 * | Revvity  | NEX446 | N/A                                    |
| [125][Tyr11]-Somatostatin-14 * | Revvity  | NEX389 | N/A                                    |

<sup>\*</sup> These are the same radiolabelled molecules, but provided in different buffers. Data shown herein were generated using NEX446. Please refer to their respective Certificates of Analysis on our WEB site for details.

Table 2. References of cell culture media and assay buffers

| Duncis              |   |
|---------------------|---|
| Provider            | Cat no  |
| Hyclone             | SH30026.02  |
| Hyclone             | SH30022.02  |
| Invitrogen          | 12634-010   |
| BioWitthaker        | 06-174G   |
| Sigma               | C8862   |
| Wisent              | 80150   |
| Wisent              | 80950   |
| Wisent              | 400-130-IG  |
| Invitrogen          | R25005  |
| invitrogen          | R210-01   |
| Wisent              | 400-160-EM  |
| GIBCO               | 14025   |
| MP Biomedicals, LLC | 101926  |
| Sigma               | A-3059  |
| Sigma               | P3143   |
| Hyclone             | SH30236.02  |
| GIBCO               | 11360   |
| GIBCO               | 25030   |
| GIBCO               | 11140   |
| Sigma               | F6886   |
|                     | Provider Hyclone Hyclone Invitrogen BioWitthaker Sigma Wisent Wisent Unvitrogen Invitrogen Wisent GIBCO MP Biomedicals, LLC Sigma Sigma Hyclone GIBCO GIBCO GIBCO GIBCO |

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