

# In vitro reverse pharmacology for characterising ligand-receptor interactions

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## **SCOPE OF THE METHOD**

| The Method relates to                               | Animal health, Human health                         |
|---|---|
| The Method is situated in                           | Basic Research, Translational - Applied<br>Research |
| Type of method                                      | In vitro - Ex vivo                                  |
| Species from which cells/tissues/organs are derived | Mammalian cell lines for heterologous<br>expression |
| Type of cells/tissues/organs                        | Mammalian cell lines for heterologous<br>expression |

## DESCRIPTION

#### Method keywords

reverse pharmacology GPCR deorphanization ligand-receptor screening cell culture

### Scientific area keywords

pharmacology neurobiology signal transduction GPCR signaling

## Method description

Reverse pharmacology is a high-throughput *in vitro* method to characterise ligandreceptor interactions. In this method, a receptor of interest is expressed in a heterologous cell line and used as a hook to fish out its ligand(s) from a library of synthetic compounds. Receptor activation is measured by monitoring secondary messengers, such as the release of calcium from intracellular storage sites, using fluorescent or bioluminescent indicators. The method can be used for highthroughput screening of ligand-receptor interactions and for in depth follow-up studies characterising the potency, affinity and downstream signalling pathways of ligand-receptor couples.

#### Lab equipment

This method requires an automated liquid handling system that can simultaneously detect fluorescence and/or bioluminescent signals, e.g. a FLIPR system. It also requires standard facilities for cell culture.

#### Method status

Published in peer reviewed journal

## **PROS, CONS & FUTURE POTENTIAL**

#### Advantages

The main advantage of reverse pharmacology is its amenability for high-throughput screening, providing the ability to perform large-scale screens of ligand-receptor interactions. In addition, no prior knowledge on downstream signalling pathways is

required to monitor receptor activation.

## **REFERENCES, ASSOCIATED DOCUMENTS AND OTHER INFORMATION**

## References

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