



# A NUCLEAR RECEPTOR DIRECTED LIBRARY

## NHRCore Library

### Highlights

- High quality, ChemBridge proprietary compounds with potential activity against nuclear receptors
- Custom select compounds to meet your requirements
- Structural analogs available for SAR studies

### Introduction

The NHRCore™ Library is a computationally selected library of more than 1,500 lead-like small molecules. NHRCore compounds are part of ChemBridge's CORE Library Stock and were selected for synthesis based on similarity to 3D pharmacophore fingerprints generated from published compounds with activity against nuclear hormone receptor targets. More than 200 synthetic schemes and more than 1,400 unique Bemis-Murcko scaffolds (atomic frameworks)<sup>1</sup> are represented.

### Selection

Compounds showing activity against nuclear hormone receptor targets were extracted from public, scientific databases. 3D conformers were generated for each agonist or antagonist compound, and 3D pharmacophore fingerprints were generated from the 3D conformers. These fingerprints were then compared to the 3D pharmacophore fingerprints generated for virtual compounds based on novel scaffolds designed by ChemBridge. Virtual compounds with a high similarity to a published active fingerprint were synthesized and included in the NHRCore Library selection.

### Potential Targets

The published actives used to generate the 3D pharmacophore fingerprints showed activity against the following nuclear receptors:

- |   |                                    |
|---|------------------------------------|
| Androgen Receptor                                       | Progesterone Receptor              |
| Estrogen Receptor                                       | Retinoic Acid Receptor (RAR) alpha |
| Farnesoid X Receptor                                    | Retinoic Acid Receptor (RAR) beta  |
| Glucocorticoid Receptor                                 | Retinoic Acid Receptor (RAR) gamma |
| Liver X Receptor alpha                                  | Thyroid Hormone Receptor           |
| Liver X Receptor beta                                   | Vitamin D Receptor                 |
| Peroxisome Proliferator-Activated Receptor (PPAR) alpha |                                    |
| Peroxisome Proliferator-Activated Receptor (PPAR) gamma |                                    |
| Peroxisome Proliferator-Activated Receptor (PPAR) delta |                                    |



## Properties

NHRCore Library compounds are lead-like and have the following physiochemical and calculated property averages:

Molecular Weight Average = 345.2

Fsp3 Average = 0.5

clogP Average = 2.0

TPSA Average = 60.3

H-bond Donor Count Average = 0.4

H-bond Acceptor Count Average = 3.6

Rotatable Bond Count Average = 4.2

## Format

- Custom select compounds according to your requirements
- Delivery in micromole or mg amounts
- Available as DMSO solutions or dry
- Formatted according to your requirements in standard or acoustic qualified plates, in 96-well format racks, or in customer specified materials

## Structures

- For a file of NHRCore Library structures, please email [sales@chembridge.com](mailto:sales@chembridge.com)
- ChemBridge can provide a standard file version or a version customized to the requirements of your project



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### References

1. Bemis GW and Murcko MA. The Properties of Known Drugs. 1. Molecular Frameworks. J Med Chem. 1996 39(15) 2887-93.