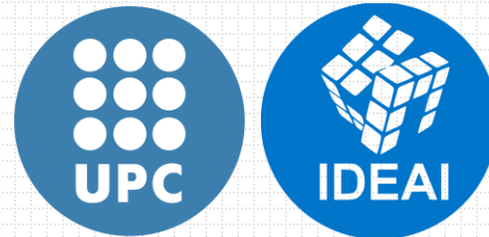


IWBIO-2022
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and Biomedical Engineering**

A Deep Learning-based method for uncovering GPCR
ligand-induced conformational states
using interpretability techniques



**Mario A. Gutiérrez-Mondragón,
Caroline König,
Alfredo Vellido.**



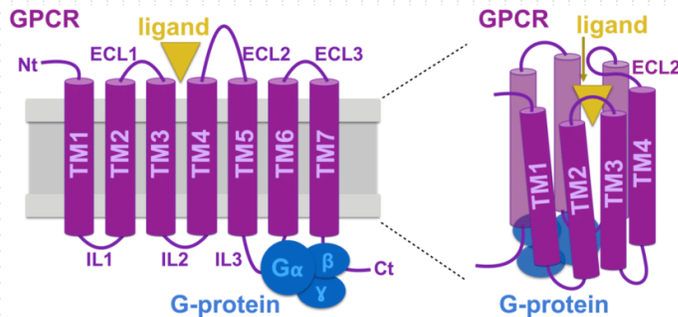
Contents

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- Materials
 - Residue Interaction Networks
 - Data pre-processing
- Experimental Setup
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- Conclusions

Study target:

G protein-coupled receptors (GPCRs).

- Primary transmembrane receptors for signal transduction.
- These proteins act as receptors of plenty of signaling molecules (ligands) - from hormones, neurotransmitters, to photons of light.
- Understanding the functional properties of these receptors is critical to deciphering the signaling process.



Schneider, J., Korshunova, K., Musiani, F., Alfonso-Prieto, M., Giorgetti, A., & Carloni, P. (2018). Predicting ligand binding poses for low-resolution membrane protein models: Perspectives from multiscale simulations. *Biochemical and biophysical research communications*, 498(2), 366-374.

Data:

Molecular Dynamic Simulations (MD).

- MD has become an established technique to explore the conformational space of proteins at an atomic level.
- MD simulations provided missing information on the dynamics of the receptors.
- The investigation of the structural information of the receptor has relevant implications in pharmacoproteomic.

Tool to perform analysis:

Deep Learning (DL)-based Models

- DL are suitable tools for knowledge discovery.
- DL models have proven to be relevant tools to alleviate and complement the challenges faced by Bioinformatics.
- However, the interpretability of the results has been poorly investigated.

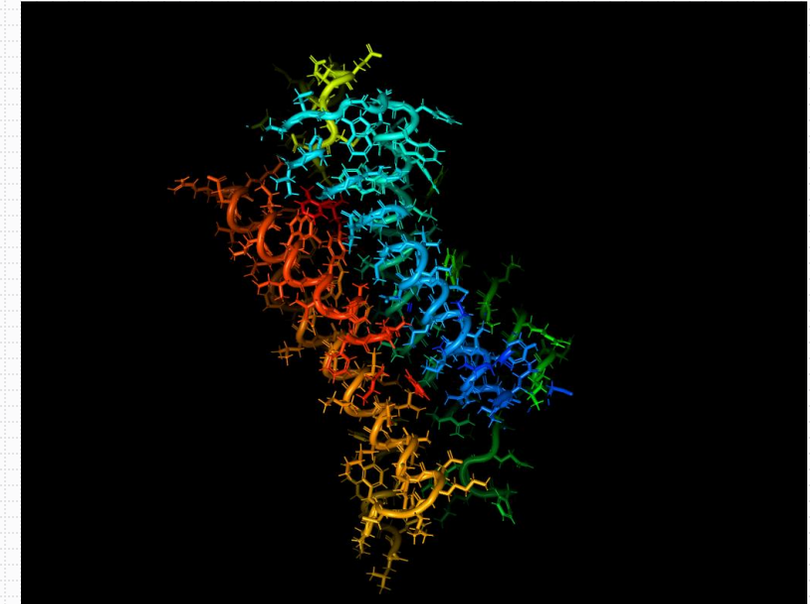
Methodology:

- Data Gathering - GPCR **β 2-adrenergic(β 2AR)** receptor with full agonist, inverse agonist and ligand-free structure.
- Data Transformation to Residue Interaction Networks.
- Model proposed: Convolution Neural Network on a Supervised classification problem on agonist-specific responses.
- Motif (residues or groups of residues) identification that induce ligand-dependent conformations through interpretability algorithms. In particular, Layer-Wise Propagation Relevance algorithm.

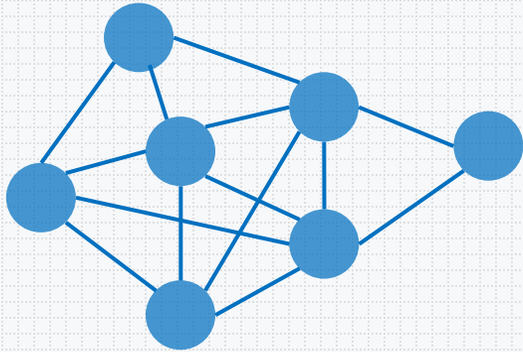
GPCR- β 2-Adrenergic Receptor cloud-based MD simulations on Google Exacycle.

Structure	Description	Type	Simulations	Length
β 2ar2RH1-a		apo (no ligand)	10,000	~12.5 ns
β 2ar2RH1-b	β 2ar2-inactive	full agonist (BI-167107)		
β 2ar2RH1-c		partial inverse agonist (carazolol)		
β 2ar2RH1-icl3		icl3 modeled - apo (no ligand)		
β 2ar3pg0-a		apo (no ligand)		
β 2ar3pg0-b	B2ar2-active	full agonist (BI-167107)		
β 2ar3pg0-c		partial inverse agonist (carazolol)		
a2a3eml	B2ar2-active	apo (no ligand)		
B1ar2y02		apo (no ligand)		

Study Case



Kohlhoff, K. J., Shukla, D., Lawrenz, M., Bowman, G. R., Konerding, D. E., Belov, D., Altman, R.B., Pande, V. S.: **Cloud-based simulations on Google Exacycle reveal ligand modulation of GPCR activation pathways.** Nature chemistry, 6(1), 15-21 (2014).

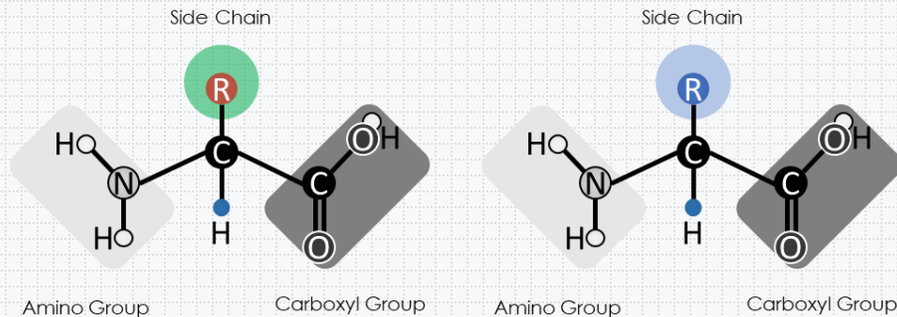


Brinda, K., Vishveshwara, S.: A network representation of protein structures: implications for protein stability. Biophysical journal 89(6), 4159–4170 (2005)

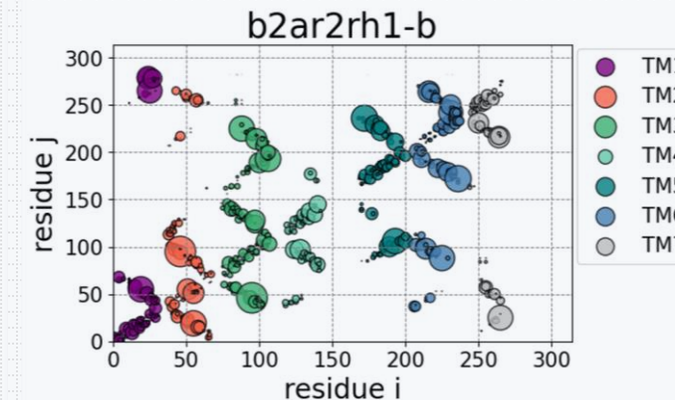
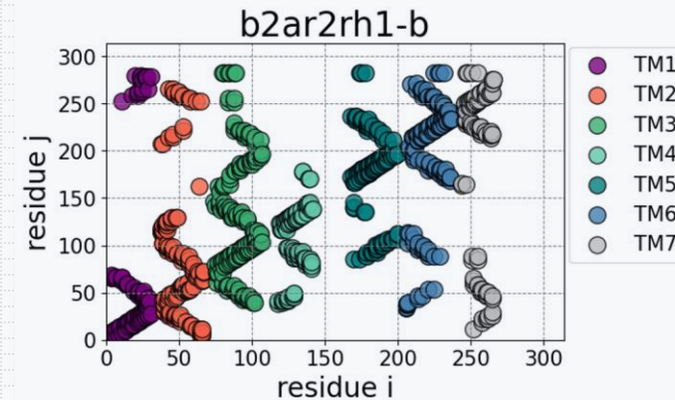
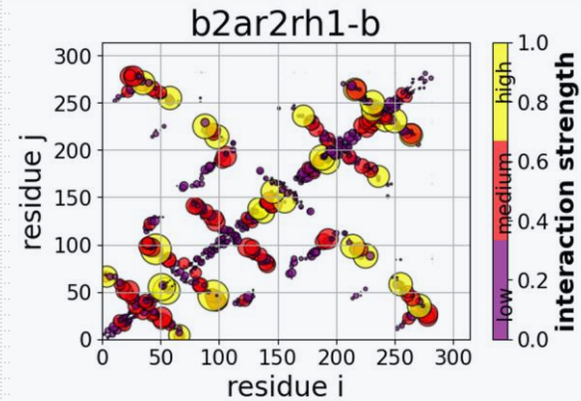
- Network representation of the protein structure to facilitate the study and analysis.
- It represent the three dimensional structure into a two-dimensional space.
- Each amino acid (residue) refers to a node, and the strength of the noncovalent interactions between two amino acids is evaluated for edge determination.

- The interaction strength is evaluated as a percentage:

$$I_{ij} = \frac{n_{ij}}{\sqrt{N_{ij}}} \times 100 \rightarrow \text{edge if } i \neq j \text{ and } I_{ij} > I_{min}$$



- n_{ij} refers to the number of distinct atom pairs between the side chains of the residues i and j , which come within a cutoff distance (4.5 Å as a default).
- N_i and N_j represent normalization factors.

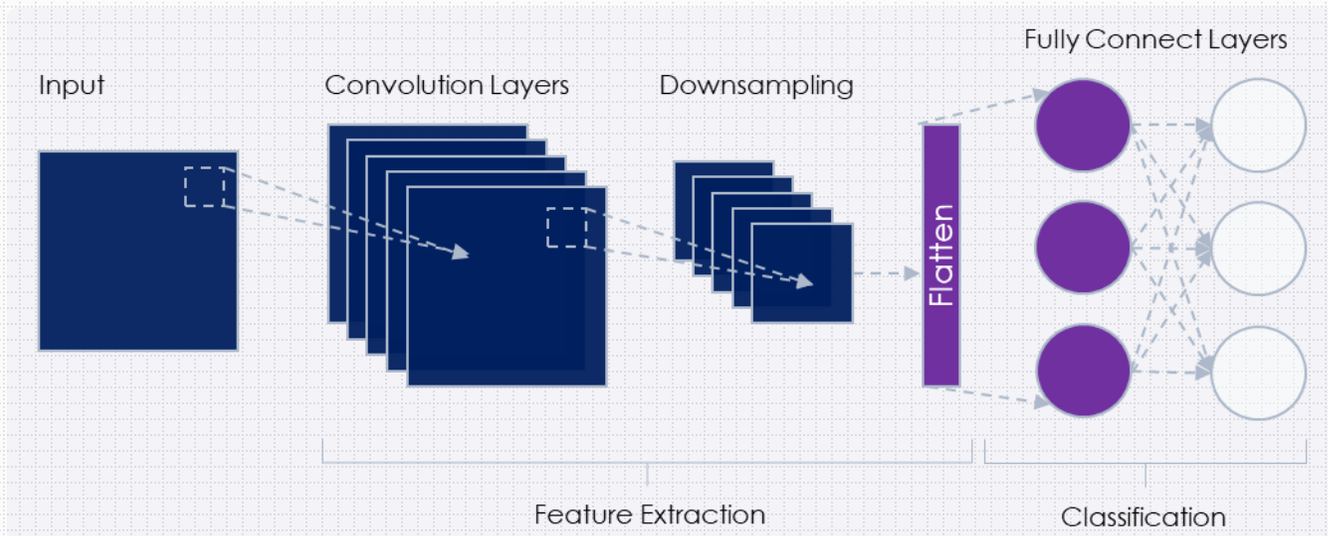


- Computations were made using PSN module from Wordom Software.

- Data Splits.

class	description	# Training samples	# Test samples
2RH1-b	β 2AR- full agonist	8,089	5,393
2RH1-c	β 2AR- inverse agonist	7,711	5,141
2RH1-icl3	apo (no ligand)	7,659	5,106
Total:		23,459	15,640

- Model Architecture Proposed.



- Overview on Convolution Neural Networks.

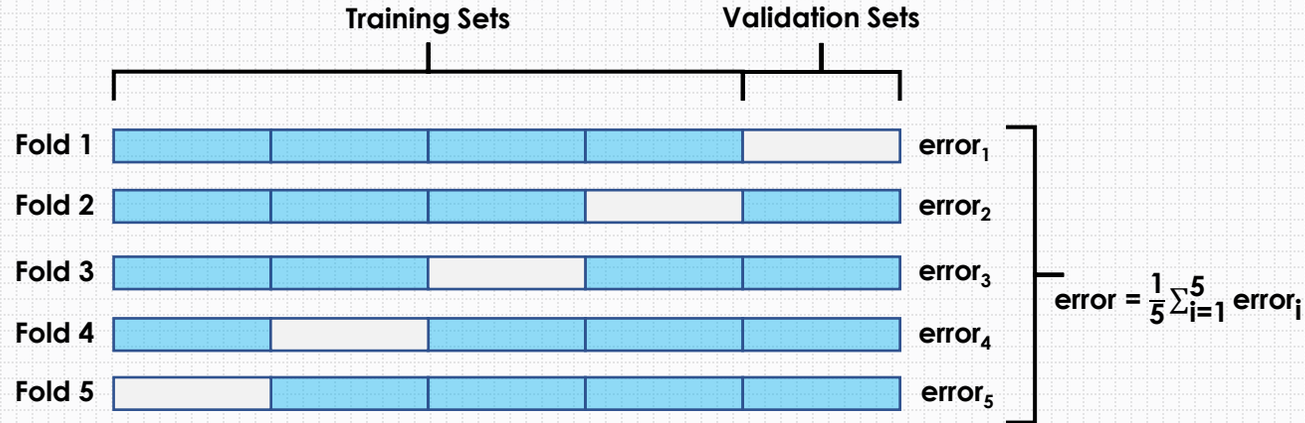
Layer(type)	Output Shape	# Parameters
Conv2d-1	[-1, 32, 310, 310]	832
ReLU-2	[-1, 32, 310, 310]	0
MaxPool2d-3	[-1, 32, 155, 155]	0
Conv2d-4	[-1, 32, 151, 151]	25,632
ReLU-5	[-1, 32, 150, 150]	0
MaxPool2d-6	[-1, 32, 75, 75]	0
Flatten-7	[-1, 18,000]	0
Linear-8	[-1, 32]	5,760,032
ReLU-9	[-1, 32]	0
Dropout-10	[-1, 32]	0
Linear-11	[-1, 3]	99

Total params: 5,768,595

Trainable params: 5,786,59

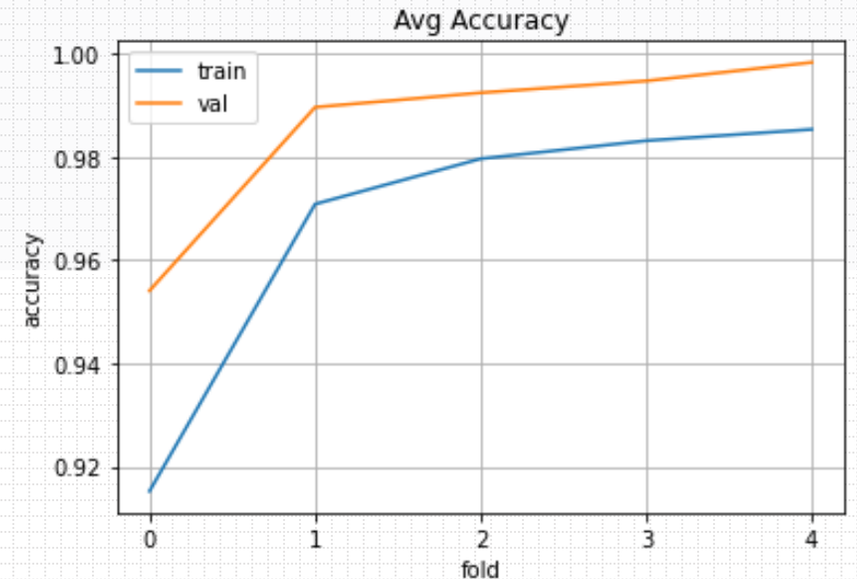
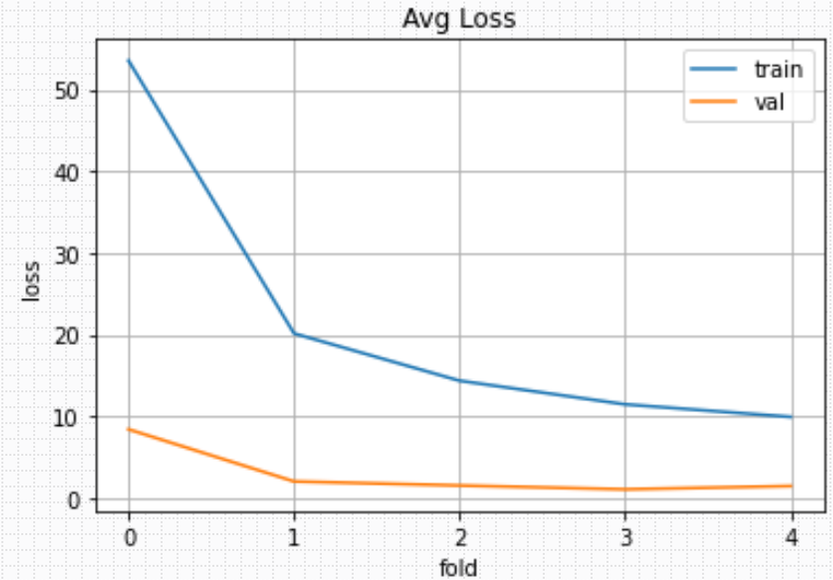
*Algorithms and computations were developed using PyTorch and Python.

- Cross K-Fold Validation Training



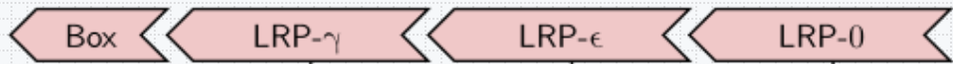
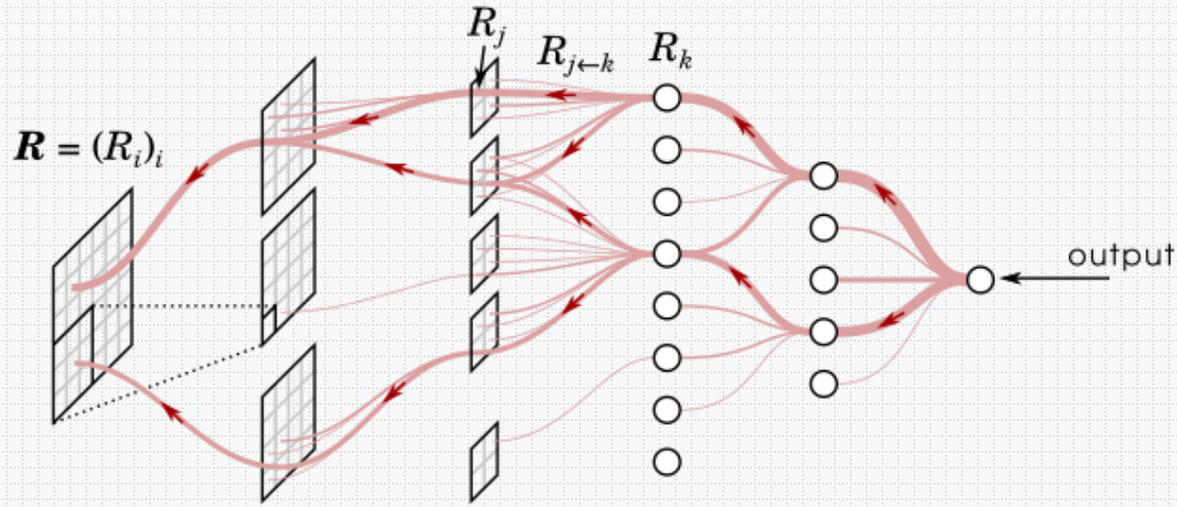
# fold	AVG Training Loss	AVG Validation Loss	AVG Training Accuracy	AVG Validation Accuracy
1	53.5658	8.3702	0.9152	0.9541
2	20.1192	1.9798	0.9709	0.9897
3	14.3261	1.4840	0.9797	0.9925
4	11.4291	1.0138	0.9832	0.9948
5	9.8927	1.4139	0.9854	0.9984
Fold AVG:	21.8665	2.8523	0.9668	0.9859

- Learning Curves



- Layer-Wise Relevance Propagation

<http://www.heatmapping.org/>



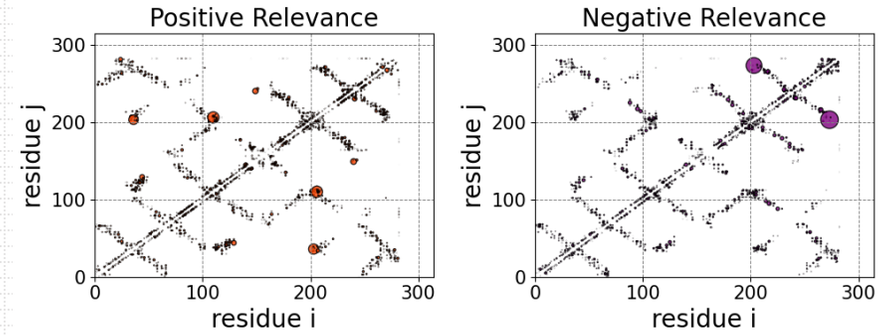
$$R_j = \sum_k \frac{a_j w_{jk}}{\sum_{0,j} a_j w_{jk}} R_k$$

$$R_j = \sum_k \frac{a_j w_{jk}}{\epsilon + \sum_{0,j} a_j w_{jk}} R_k$$

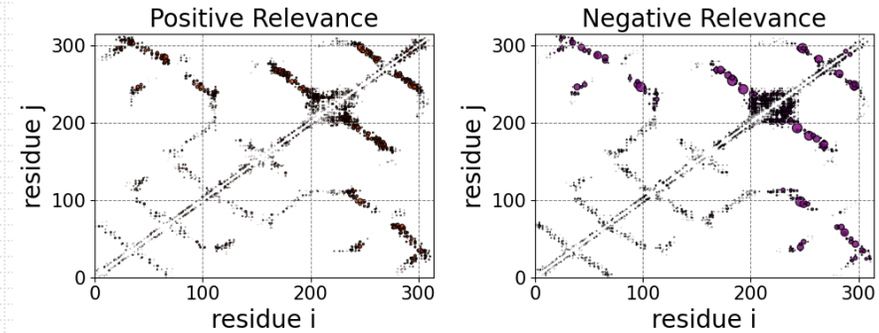
$$R_j = \sum_k \frac{a_j (w_{jk} + \gamma w_{jk}^+)}{\sum_{0,j} a_j (w_{jk} + \gamma w_{jk}^+)} R_k$$

Montavon, G., Binder, A., Lapuschkin, S., Samek, W., & Müller, K. R. (2019). Layer-wise relevance propagation: an overview. *Explainable AI: interpreting, explaining and visualizing deep learning*, 193-209.

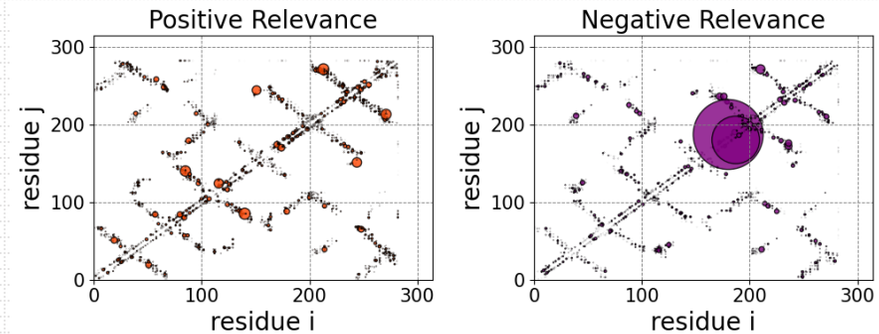
a) AVG Relevance $\beta 2ar2$ with full agonist.



b) AVG Relevance $\beta 2ar2$ with inverse agonist.



c) AVG Relevance $\beta 2ar2$ free ligand.



- Motifs Identification

- Relevance Per Transmembrane.

a) Transmembrane AVG Relevance β 2ar2 with full agonist.

region	# RI	AVG R
TM1	178	0.002775
TM2	219	0.003977
TM3	236	0.003365
TM4	126	0.005040
TM5	211	0.004188
TM6	269	0.005906
TM7	186	0.003853

b) Transmembrane AVG Relevance β 2ar2 with inverse agonist.

region	# RI	AVG R
TM1	183	0.006238
TM2	239	0.007591
TM3	260	0.013146
TM4	116	0.018101
TM5	219	0.011499
TM6	291	0.011066
TM7	213	0.012098

c) Transmembrane AVG Relevance β 2ar2 free ligand.

region	# RI	AVG R
TM1	214	0.005606
TM2	179	0.002913
TM3	251	0.002294
TM4	164	0.002551
TM5	445	0.003197
TM6	257	0.012646
TM7	66	0.002494

- The study of the conformational space of GPCRs has critical implications in Drug Discovery process.
- Molecular Dynamics and Machine Learning-based models enable a further understanding of molecular processes providing relevant insights into the dynamics of proteins.
- The proposed methodology addresses the transformation of the raw simulations into a Protein Interaction Network.
- We have presented a Convolution Neural Network with high accuracy on a classification problem to provide evidence of the ligand-specific GPCR activity.
- We also include an interpretability study of the prediction results. Therefore, we can identify relevant features (motifs) that underlie conformational rearrangements influenced by ligands.
- Importantly, we provide the trustworthiness of the proposed model and a method to further assess its predictions in this domain.

Thank You!

mario.alberto.gutierrez@upc.edu
caroline.leonore.konig@upc.edu
avellido@cs.upc.edu