Binding site identification of G protein-coupled receptors through a 3D Zernike polynomials-based method: application to C. elegans olfactory receptors

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Received: 15 June 2021 / Accepted: 18 November 2021 / Published online: 1 January 2022
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Abstract
Studying the binding processes of G protein-coupled receptors (GPCRs) proteins is of particular interest both to better understand the molecular mechanisms that regulate the signaling between the extracellular and intracellular environment and for drug design purposes. In this study, we propose a new computational approach for the identification of the binding site for a specific ligand on a GPCR. The method is based on the Zernike polynomials and performs the ligand-GPCR association through a shape complementarity analysis of the local molecular surfaces. The method is parameter-free and it can distinguish, working on hundreds of experimentally GPCR-ligand complexes, binding pockets from randomly sampled regions on the receptor surface, obtaining an Area Under ROC curve of 0.77. Given its importance both as a model organism and in terms of applications, we thus investigated the olfactory receptors of the C. elegans, building a list of associations between 21 GPCRs belonging to its olfactory neurons and a set of possible ligands. Thus, we can not only carry out rapid and efficient screenings of drugs proposed for GPCRs, key targets in many pathologies, but also we laid the groundwork for computational mutagenesis processes, aimed at increasing or decreasing the binding affinity between ligands and receptors.

Keywords GPCR · Protein-ligand interaction · Binding site prediction · Protein structure

Introduction

The identification and characterization of ligand binding sites in proteins is a fundamental step for structure-based drug design [1, 2]. Among all protein families, G protein-coupled receptors (GPCRs) are probably the most popular family of drug targets, accounting for about 34% of all pharmaceuticals approved by the US Food and Drug Administration [3].

GPCRs play a key function in many biological processes, mediating a large number of cellular responses to external stimuli, such as light, odors, hormones, and growth factors. The communication between the outside and the inside of the cell takes place through the interaction between GPCRs and ligands in the extracellular environment, allowing the protein to bind, in an intracellular environment, with a heterotrimeric G protein [6]. This process is followed by various forms of signal transduction, all these events originated by the interaction of the GPCR with a specific ligand [7].

Therefore, the understanding of the interaction mechanism between small molecules and GPCRs represents a fundamental step for the design of new pharmaceutical compounds [8]. Among all the GPCR-ligand interactions, a relevant part is the recognition mechanisms between odors and their corresponding Odor Receptors (ORs), which...
represent the largest subfamily within the GPCRs family and are present in all multicellular organisms [9–11]. Indeed, these interactions are fundamental for many biological applications [1]. For instance, the level of expression of ORs in tumoral tissues is noticeably different compared to healthy tissues, and, currently, these receptors are candidates as new targets for diagnosis and therapeutics, but a complete comprehension of their biological role in cancer is still elusive [12]. More specifically, it is still unclear how olfactory comprehension of their biological role in cancer is still elusive [13–15], making the design of specific receptors for any given odorant difficult [16]. Di Pizio et al. recently reviewed in detail the role of ORs in physiological and pathological processes [17].

As easily predictable given the importance of GPCR proteins, in recent years many biochemistry and bioinformatics approaches have been developed to study drug-receptor binding [7, 18, 19], and some of them have been even applied for the prediction of their corresponding binding affinity [20, 21]. As for other protein-ligand prediction methods, most of them are based on machine learning methods such as SVM (Support Vector Machine) [22–24] and Neural Network [25, 26], but unfortunately only a limited amount of structural data are available for training these models.

Indeed, since GPCRs are integral membrane proteins containing seven transmembranes (TM) α-helices and are characterized by flexible and dynamic structures, it is very difficult to obtain experimental structures based on biochemical and crystallographic experiments [27]. For this reason, computational studies aimed at modeling the three-dimensional structure and, eventually, the association between GPCR and ligand are fundamental to investigate the GPCR-ligand interaction properties [28].

Here we present a new procedure based on a parameter-free computational approach, able to recognize, completely unsupervised, if a region of a GPCR protein can be involved in an interaction with a given ligand.

The method is based on the 3D formalism of the Zernike polynomials [29, 30] and it can characterize with an ordered set of numerical descriptors the morphological properties of a molecular surface. In particular, we computed the molecular surface [31] of a protein and we extract the portion generated by a specific set of residues (patch). This patch can be represented as a function in the 3d space that can be expanded on the basis of the Zernike polynomials (see Methods, Eq. 3). The coefficients of such expansion are determined by the form of the specific patch analyzed since they represent the weights that each polynomial assumes in the expansion. The Zernike descriptors are obtained by computing the norms of such coefficients, and they are invariant under rotation and translation in 3D space. Therefore, each selected molecular shape is summarized with its corresponding Zernike descriptors. We calculated the Zernike descriptors for the ligand binding site and for the ligand itself, being able to compute their compatibility simply by calculating the distance between the corresponding descriptors (see Figure 1).

This framework allows us to effectively evaluate the compatibility between two molecules because, in principle, two perfectly fitting surfaces are identical and then they share the same Zernike descriptors. Therefore, the distance observed between interacting surfaces (ligand–binding site) is significantly lower than the one observed between non-interacting surfaces (ligand – randomly selected exposed region of the same size of the binding site). In the last years, the Zernike formalism has been widely applied for the study of molecular complementarity [32–38].

Thus, we collected a large structural crystallographic dataset of GPCR-ligand experimental complexes. Computing the shape complementarity employing Zernike descriptors, the difference, in terms of ligand compatibility, between binding sites or randomly surface regions is clearly recognizable.

We thus applied the proposed strategy to investigate the ligand-receptor interaction in the nematode Caenorhabditis elegans. This tiny roundworm detects a large class of volatile odorants, making the olfaction its primary sense, and represents a model organism for studying and addressing the biology of olfaction. The olfactory system of C. elegans comprises approximately 32 chemosensory neurons and more than 5% of its overall genes are related to sensing chemical cues in the environment [39]. Unlike mammalian olfactory neurons, each expressing just one kind of olfactory receptor, C. elegans chemosensory neurons express more than one receptor per cell and each worm’s chemosensory neuron may respond to a large variety of molecules. As a consequence, despite the low number of constituting cells, the olfactory circuit of C. elegans recognizes hundreds of molecules at a wide range of concentrations [13, 39, 40].

Studying the olfactory network in C. elegans may help in addressing a challenge more demanding in higher brain systems: how does the neural coding strategy of odors actually work? Despite the compact nervous system, the nematode is capable to sense and integrate into reproducible behaviors a multitude of olfactory signals. C. elegans provides therefore a unique and easy-to-manage platform to study neural circuits underlying smell.

However, the large co-expression of GPCRs within olfactory neurons makes genetic screens ineffective in linking receptors to odors and only a few GPCRs have been linked to the corresponding ligands [41]. Using the method presented in this paper, we build a comprehensive list of odorant-receptor possible associations in an olfactory neuron of C. elegans. However, it is worth noting that this protocol can be applied in analogous types of analysis for every kind of chemosensory receptors.
Specifically, we select 21 among GPCRs expressed on the main olfactory neuron mediating attraction in *C. elegans*, the AWC neuron [42]. AWC neurons are a pair of olfactory neurons critical for chemotaxis to volatile odors (e.g. attraction to benzaldehyde, butanone, isoamyl-alcohol, 2,3 pentanedione, and 2,4,5 trimethylthiazole) [13]. Their activation induces local search behavior and promotes turns. We tested these 21 GPCRs against 18 putative ligands we identified in a recent paper using a sequence-based methodology [43]. In particular, sampling the 21 olfactory receptors of *C. elegans*, we computed the shape complementarity between each receptor patch and the proposed ligands. Thus we can evaluate these results with respect to the typical complementary values observed in experimental GPCR-molecule interactions. From this perspective, with an *in-silico* characterization we propose a set of GPCR-ligand interactions in *C. elegans* since our formalism shows that that couple exhibits an exceptionally high shape complementarity.

**Results and discussions**

The identification of the binding region responsible for the capture of a chemical compound on a GPCR is a key aspect both to understanding the molecular recognition mechanism and to open the door to new methodologies for drug design and optimization of ligand-GPCR interactions. For example, it is possible to design an *ad-hoc* compound able to compete with the physiological ligand or, employing computational mutagenesis protocols, amino acid substitutions can be carried out to increase (or decrease) protein-ligand binding compatibility.

In this study we developed a new computational procedure, completely based on shape complementarity analysis, able to distinguish the specific interaction occurring between a ligand and its GPCR binding region from the non-specific ones, involving a set of randomly sampled regions on the surface of the protein itself. To this end,
we selected a large experimental dataset and analyzed the geometric properties of the molecular surface regions by adopting the Zernike polynomials description, which is a powerful tool in capturing the structural determinants of both protein-protein and protein-ligand recognition [32–36, 38, 44].

After demonstrating that the sensibility of the employed representation is good enough to properly work even when a predicted protein structure is considered (a significant aspect due to the lack of the experimentally resolved GPCRs structure), we applied the computational procedure to investigate the odor-receptor recognition mechanisms of the model organism *C. elegans*. Here, we propose some possible associations between the putative ligands identified in [43] and a set of 21 GPCRs belonging to AWC neuron of *C. elegans*.

### Detection of ligand-specific binding site regions on GPCRs

We collected a dataset of 287 experimentally solved GPCR-ligand complexes. For each structure, we defined the GPCRs binding sites (BS) as the set of residues whose *Cα* are closer than 6 Å to any atoms of the ligand.

We first analyzed the frequencies of amino acids occurrences in the binding regions, as shown in the barplot in Fig. 2A. Comparing our results with the corresponding ones regarding the binding sites of a systematic dataset of protein-ligand complexes (about 4000), some remarkable differences turn out [45]. Some non-polar residues, such as Ala and Val, increase largely their frequencies, while charged residues (Arg, Lys, Glu, Asp) are very low represented in GPCRs binding sites. Moreover, it can be noted that interactions mediated by Tyr and Trp aromatic residues are less common.

![Fig. 2](image)

Fig. 2 Analysis of the interaction between ligands and GPCRs binding sites: application of the Zernike formalism. A The amino acid distribution regarding the amino acids involved in binding. B ROC curves obtained using the Zernike Descriptors. The green line (AUC = 0.77) regards all the dataset, the red line (AUC = 0.60) consider only the interaction between GPCR protein and peptide ligand, while the blue line is related to the ligands with a molecular weight lower than 500 Da (AUC = 0.81). The molecular images represent an example of GPCR-small molecule and GPCR-peptide recognition, respectively on the left and the right. C The same as in A, but the results are grouped according to the membership of each GPCR-ligand complex to one of the quartiles of Zernike accuracy (Cyan = 1st quartile means high complementarity while Brown = 4th quartile means low complementarity.) D The number of Binding site residues as a function of the molecular weight of the ligand. The points are colored according to the membership to one of the quartiles of Zernike accuracy (Cyan = 1st quartile means high complementarity while Brown = 4th quartile means low complementarity.)
than in the general case: on the contrary, some polar residues (Thr, Ser) increase their occurrences. Taken together, these results seem to give insights into the peculiar nature of the GPCRs binding sites, often located in a very internal area with limited solvent exposure.

We defined on each GPCR a set of randomly selected surface regions (Binding Site Decoy, BSD), in such a way that these are as comparable as possible with actual BS (See Methods Section). Thus, we calculated the Zernike descriptors of both the BS and the BSD, and the ligand as well. We, therefore, summarized in an ordered vector of numerical values the geometric properties of the protein patches (BS and BSD) and the ligand. Since the shape of two perfectly fitting surfaces is the same, we quantified the complementarities between protein and ligand computing the distance between the corresponding Zernike descriptors, where lower is the distance higher is the complementarity (See Methods section).

This analysis aims to verify if the Zernike description can distinguish, in terms of shape complementarity, specific interactions (ligand-binding site) from non-specific ones (ligand-decoys), analyzing in this way both the performance of the designed method and the role of shape complementarity in receptor-ligand recognition. The main hypothesis of this study is that the binding between ligand and receptor is mainly mediated by the geometric complementarity of the two molecules. In this scenario, since the actual binding sites should achieve a higher shape complementarity with the ligand than other surface regions, the distance between the descriptors of the ligand and the binding site should be lower than the ones involving the decoys.

Collecting the results regarding all the complexes in the dataset, the distribution of the specific interaction Z-score is characterized by values significantly lower than 0, meaning that the formalism on average capture the differences in shape complementarity between specific and non-specific interactions. In particular, the specific interaction Z-scores are characterized by a mean of -1.07 and a standard deviation of 1.18.

In the light of these results, we performed a ROC analysis to quantify the protocol binding site prediction power (Fig. 2B): labeling as positive the actual BS and as negatives the BSD, and using the Zernike distances with the cognate ligands as discriminant, we obtained an area under the curve (AUC) of 0.77, meaning that we satisfactorily spot the correct protein region-ligand associations.

Interestingly, our method shows different performances depending on the kind of ligand bound by the GPCR. Indeed, when only peptidic ligands are considered the performances drastically decrease (AUC = 0.60), while when only small ligands (molecular weight lower than 500 Da) are examined, the computational classification protocol works even better than in general case (AUC = 0.81). Since peptides are usually characterized by a high molecular weight in respect to non-peptidic ligands, a possible interpretation can be that ligand-pocket shape complementarity is sufficient for the identification of pockets suitable for the recruitment of small compounds, while for the recognition of heavier ligands it is also necessary a good chemical-physical interaction [46, 47]. Indeed, as shown in the molecular images in Fig. 2B, the shape complementarity can be caught by Zernike description very effectively when a small molecule is considered, since a small pocket is perfectly built around the ligand. When the molecule recognized by a GPCR is characterized by a high molecular weight (as the peptides usually are), the binding site that recognizes it is connected to the outside solvent. Likewise, a large portion of the ligand molecular surface is not optimized to bind the protein binding site since it is not in direct contact with the protein. Therefore, when such molecule is expanded on the Zernike basis, the descriptors will represent also the part of the surface that is not complementary to the protein binding site, resulting inevitably in a decrease of protein-ligand shape complementarity.

To verify if certain residues can mediate preferentially shape complementarity with ligands, we thus grouped GPCR-ligand complexes into 4 categories depending on the accuracy the methods achieve in detecting the actual binding site. In particular, each GPCR was assigned in the corresponding quartile of the Z-scores distribution, so as the 1st quartile contains the complexes best classified while the 4th one incorporates the GPCR-ligand complexes with the worst scores.

In Fig. 2.C we reported the frequencies of amino acids in the binding site separated according to the 4 quartiles. Interestingly, in the GPCRs binding sites characterized by a high shape complementarity are over-expressed some amino acids, for instance Valine (Val) and Serine (Ser), suggesting that such small and flexible residues can effectively adapt themselves to accommodate the ligand.

Moreover, in Fig. 2.D, we showed the number of residues involved in binding sites as a function of the ligand molecular weight. As expected, a linear correlation between two quantities exists (Pearson correlation coefficient of 0.32, p value < 10^-5). The points in this plot are colored in accordance to the quartile membership of each molecular complex: even if lighter ligands are slightly better predicted (see Figure 2.B), there are no significant separations between these groups and therefore we can conclude that the description based on Zernike moments, in this range, is not biased by the binding site size (note that in this analysis we not consider peptide ligands).

**Evaluation of ligand flexibility in shape complementarity with GPCRs**

The compatibility between a protein binding site and a ligand is dependent on the conformation adopted by the
ligand. Bound conformations are usually characterized by a shape complementarity higher than the one observed when unbound states (APO conformation) are investigated. Thus, in this section, we investigated how ligand flexibility impacts the compatibility with the GPCR binding site. In particular, we selected a subset of ligands and we performed a molecular dynamics analysis to establish how their flexibility impacts the compatibility with the protein binding site. We summarized the results of this analysis in Fig. 3.

Looking at the distribution of the molecular weights of the ligands in our dataset, we selected one ligand from each decile. Therefore, we constituted a subset of 10 ligands stratified in terms of molecular weight (see Fig. 3A). To explore the conformational space of each ligand, we extracted 100 different ligand conformations sampling uniformly each ns a 100-ns long molecular dynamics simulation. Finally, we summarized the shape features of each obtained conformation by computing its Zernike descriptors. For each simulation, we computed
the Zernike descriptors distances between the experimental ligand conformations and all the molecular dynamics frames (Fig. 3B blue boxplots). Moreover, we computed the distances between all the explored frames in the simulation (Fig. 3B red boxplots). Therefore, the red distributions represent the level of variability of each structure. The blue distribution, the distances between all the possible conformations with the experimental one, is typically higher than the red one. This is because the bound state represents a small region in the configurations space, often not located in a very explored area. This notwithstanding, the overlapping between the 2 distributions means that the molecular dynamics of the ligand explore some conformations similar to the bound state.

We studied as the difference between the unbound states and the bound one, in terms of Zernike descriptors, is correlated with the pocket-ligand specificity (Fig. 3C). In particular, for each frame of the simulations, we calculated the complementarity between the particular ligand conformation and the experimental binding pocket or the decoys. Thus, we normalized with the Z-score as explained in the manuscript, so as the Z-score of the ligand-binding pocket association is a measure of the sensibility of our approach (the more negative the Z-score is, the more correct is the association between ligand structure and the pocket). As evident, exist a correlation between the difference with the bound state and the capability of recognizing the binding pocket (Pearson correlation coefficient = 0.45). It is interesting to note that the points out of the linear trend are mainly belonging to the dynamics of K86(5xpr), a ligand that we can not correctly assign to its pocket even using the experimental conformation (its experimental Z-score is 1.55).

Finally, we calculated the area under the ROC curve obtained when molecular dynamics frames are considered (Fig. 3D). Ordering the frames according to their similarity with the experimental ligand structure, we report in the figure the mean AUC we obtained considering frames from most to least similar (blue curve) or from least to most similar (red curve). The orange dashed line represents the AUC we obtained with experimental conformation, for this specific subset of GPCR-ligand interactions. It has to be noted that when we consider the structure most similar to the experimental one, the mean performance we get (AUC = 0.75) is very similar to the experimental case (0.76), while the mean performance obtained when the ligand structure is least similar to the bound state is significantly lower (0.67). Including a number progressively higher of structures, by construction, the 2 performances tend to become more and more similar until the perfect identity of the last point (when we consider all the structures of the simulations in both cases). Intuitively, as long as the unbound structure is similar to the bound one, we obtain performances very similar to those obtained in such case; indeed, the conformations explored in dynamics are close enough to the bound state to make possible a satisfying performance.

### The application of Zernike formalism to GPCRs models

In the previous section, we demonstrated that our approach based on Zernike formalism obtains very good results in binding site identification when applied to experimental GPCR-ligand structures. Unfortunately, the structural details of a very large fraction of GPCRs are still lacking [48, 49], also because of the inherent difficulties of working with membrane proteins.

Therefore, to extend the applicability of this formalism to the whole GPCR family, we need to establish if the shape representation employing Zernike descriptors works properly even when we deal with predictions of GPCR structures provided that, obviously, the results of this study will depend on the quality of the predicted structures.

In light of these considerations, we tested our protocol selecting a subset of the structural dataset and investigating the similarities between predicted and experimentally solved structures. In particular, choosing 14 structures (approximately 5% of the entire dataset) belonging to different classes of GPCRs (according to GPCR-EXP database [50]), we extracted the protein amino acid sequence and we modeled the structure using the I-Tasser web-server [51], excluding from the possible template for homology modeling known structures with too high sequence similarity (all possible templates with sequence identity higher than 25% have been excluded).

We thus compare the native experimental structures and the predicted ones. For each GPCR, we defined for both the experimental and predicted structures a set of surface regions, basically centering on each residue a spherical patch with a radius of 9 Å. We computed the Zernike descriptors of all these patches. It is hence possible to define on one hand the specific similarities, the distances between the corresponding patches on the experimental and modeled structure (those centered on the same residues), and on the other hand the non-specific similarities, the distances between all the couples of non-corresponding patches (centered on different residues), as illustrated in Fig. 4.A. If the modeling works properly, the mean of the specific similarities has to be lower than the mean of the non-specific similarities.

Therefore, for each protein sampled with \( i = 1,...,N \) patches, it holds

\[
S_s = \frac{1}{N} \sum_{i=1}^{N} S_{xi} = \frac{1}{N} \sum_{i=1}^{N} \text{dist}(E_i, M_i) \tag{1}
\]

\[
S_{ns} = \frac{2}{N(N-1)} \sum_{i=1}^{N} \sum_{j>i}^{N} S_{xij} = \sum_{i=1}^{N} \sum_{j>i}^{N} \text{dist}(E_i, M_j) \tag{2}
\]
where $E_i$ and $M_i$ are the Zernike descriptors of the $i$-th patch relative to the Experimental or Modeled structures, respectively. $S_s$ and $S_{ns}$ thus represent the specific and non-specific similarity relative to the examined protein. We therefore defined the Zernike Sensitivity as $S_Z = S_{ns} - S_s$, meaning that when $S_Z > 0$ our representation preferentially associate corresponding regions.

In Fig. 4.B, we reported for each protein the Zernike Sensitivity as a function of the Root Mean Square Deviation (RMSD) between the modeled and the experimental structure (both the superposition and the calculation was based on $Ca$ atoms). When the model has a good quality (low RMSD) the Zernike protocol performs well in recognizing the same region in the 2 structures (high Zernike sensitivity) (Pearson Correlation Coefficient $R = -0.65$, p-value $= 0.01$).

Moreover, beyond the global quality of a model and the corresponding Zernike accuracy, it is interesting to study even the local goodness of the predicted structure. Indeed, the I-Tasser web server returns an estimation of the exactness of the single residue position prediction (L-score [52]).

Calculating the mean L-score of each region previously defined we basically defined the confidence of this region prediction. For each experimental-modeled pair of structures, we normalized the distribution of specific similarities and non-specific similarities with the Z-score, similarly to what was done above. In this way, when a region exhibits a
low similarity Z-score it means that the specific similarity is better than the typical non-specific one.

In conclusion, we compared the similarity Z-scores with the mean L-score of the region and we reported the results in the boxplot shown in Fig. 4:C: as expected, when the local predicted quality of the model is good (low L-score) even the Zernike formalism can recognize better the similarity between predicted and experimental structure.

In this section, we demonstrated that both from the global or local point of view, as long as the protein modeling reaches good results the Zernike description grasp the similarity of the regions and can be therefore applied, with good approximation, on predicted structures.

Olfactory receptors: the case of Caenorhabditis elegans

In this section we focused on the olfactory receptors of the model organism C. elegans [42].

In a recent work [43], we have designed a new sequence-based strategy to determine the association between each OR and a set of new possible volatile compounds, based on the hypothesis that similar compounds bind similar proteins [54]. In that work, we produced a list of possible ligands for C. elegans GPCRs, reported in Table 1, where compounds have been considered if they have been found experimentally in complex with a GPCR at least once.

To structurally predict possible associations between ligands and receptors, in this section we assessed the shape complementarity observed between each protein and each ligand.

Since the structural details of these proteins are still not available, we modeled them using GPCR I-Tasser web-server. In the previous section, we have shown that when the model is good enough our approach can accurately summarize local shape information. Here, since we can not make the comparison with the experimental structures, we collected for each model the cognate C-score, a confidence score for estimating the quality of the predicted model, as returned by the web-server. As stated here [55], since the C-score highly correlates with the RMSD, when it is higher than −1.5 the model can be considered satisfactory. In panels A, B, C of Fig. 5, the GPCRs related to models with a C-score higher than −1.5 are highlighted in blue print while those with a C-score lower than −1.5 are highlighted in red print.

Thus, we extracted from each GPCR a set of possible ligand binding sites, simply centering on a residue and selecting all the residues closer than 9 Å to it. Selecting the surface points generated by these residues, we defined the patches and computed the Zernike descriptors of each of them. Moreover, we described with the corresponding Zernike descriptors also all the 18 chosen ligands.

Table 1 The putative GPCRs ligands selected in this analysis

| PDB entry | Ligand       | Pubchem CID [53] | X-ray resol [Å] |
|-----------|--------------|------------------|-----------------|
| 5U09      | Taranabant  | 11226090         | 2.6             |
| 4N6H      | Naltrindole | 5497186          | 1.8             |
| 5NDD      | AZ8838      | 126961334        | 2.8             |
| 5ZBH      | BMS-193885  | 9960164          | 3               |
| 4IB4      | Ergotamine  | 8223             | 2.7             |
| 5G53      | N-Ethyl-5’-Carboxamido Adenosine | 8804901 | 3.4 |
| 5L7D      | Cholesterol | 5997             | 3.2             |
| 4DJH      | JDTic       | 9956146          | 2.9             |
| 5TGZ      | AM6538      | 46912833         | 2.8             |
| 1U19      | Retinal     | 638015           | 2.2             |
| 2YDO      | Adenosine   | 60961            | 3.0             |
| 4Z34      | ONO9780307  | 252166790        | 3.0             |
| 4BNV      | Cyanopindolol | 46937143        | 2.1             |
| 4JSP      | ATPgammaS   | 44123300         | 3.3             |
| 3EML      | ZM241385    | 176407           | 2.6             |
| 3UON      | Quinuclidinyl benzilate | 688566 | 3.0 |
| 2RH1      | (S)-Carazolol | 13023332      | 2.4             |
| 4IAQ      | Dihydroergotamine | 10531  | 2.8 |

The first column reports the PDB code of the experimental structure where each ligand has been found in interaction with a GPCR, and from which is taken its 3D structure.

We then computed the complementarities between ligands and GPCRs regions, labeling each ligand-GPCR pair with the higher shape (lower distance) compatibility between the compound and one of the protein pockets (Fig. 5A). At the same time, we reported in Fig. 5B the local mean L-score of the selected protein pockets, characterized by the best complementarity value of the ligand-GPCR pair.

We established an association between a ligand and a GPCR if the Zernike distance is lower than 0.09, the upper bound of the 5% lower interval of the distribution of distances between random surface regions and ligands in the structural dataset. This means that an association is formed when there is a probability lower than 5% that the observed shape complementarity belongs to the distributions of “non-associations”.

Moreover, to further control the goodness of the ligand-protein pairing, we required also that the selected binding region exhibits a good mean L-score, belonging to the first quartile of L-scores considering all the protein regions defined in our modeled GPCRs.

The proposed interactions are depicted in Fig. 5C, where a colored spot is placed when a ligand-protein association is hypothesized. It is interesting to note that 2 ligands, AZ8838 (8TZ) and Adenosine (ADN), show a high number of possible associations: this behavior can be due to the small
size of these compounds (molecular weight of 219 and 254 Da respectively), that make them compatible, in terms of shape complementarity, with a larger set of protein binding pockets.

Conclusions

Given their centrality in several biological processes and their importance as a drug target, GPCR proteins have been largely studied to understand the mechanism of interaction with ligands. Here, using a method based on Zernike polynomials on a large dataset of ligand-GPCR X-ray structures, we quantitatively demonstrated that the ligand recognition is mediated by a high shape complementarity with the protein binding sites. We thus assessed the impact of ligand flexibility, studied with molecular dynamics simulation, on the sensibility of our method: we showed that as long as the conformational exploration can reach molecular conformations similar to the experimental ones, our formalism reaches performances in line with those obtained using experimental data. Moreover, the reliability of the Zernike formalism is confirmed when predicted protein structures are used, and this aspect is particularly relevant considering the inherent difficulty in obtaining experimental GPCR structures.

Based on these results, we designed a new fully computational procedure for the GPCR-ligand association and binding sites prediction. Indeed, working on some proteins of the olfactory network in *C. elegans* and on a set of possible ligands, we can hypothesize an association if the complementarity between a small molecule and a confidently predicted protein pocket is exceptionally high.

Undoubtedly, other elements characterizing ligand-pocket compatibility have to be included in the evaluation to improve the method performance. The expansion in the Zernike polynomials can be used also in such a direction, since any Physico-chemical property, such as electrostatic potential or hydrophobicity profile computed on the molecular surface, can be represented with Zernike descriptors [33, 38, 56].

From this point of view, our results can represent a promising step toward the elusive and important goal of predicting GPCRs specificity. The development of reliable and accurate computational methods for GPCR-ligand association can have numerous applications, from driving the design of a drug able to compete for binding in a given pocket to the evaluation of which amino acid substitutions can disrupt a molecular binding.

Materials and methods

Datasets

For the study of shape complementarity between small molecule and the corresponding binding region of the
GPCR, we used GPCR-EXP database [50], selecting only GPCR-ligand experimentally solved complexes. Among all the complexes available, we have considered the complexes experimentally solved in X-ray crystallography. We have obtained a total of 287 structures and we called it “GPCR-Lig dataset”.

In order to study the molecular mechanisms of C. elegans odor recognition via receptor-small molecule interaction, we have selected a set of 21 GPCR genes, 20 of them encoding GPCRs expressed on AWC neurons: sra-13, srb-16, srd-5, srd-17, sre-4, sri-14, srj-21, srj-22, srx-3, srx-5, srx-37, srt-7, srt-28, srt-29, srt-45, srx-1, str-1, str-2, str-130, str-199 [57]. For this study we include also odr-10, a gene expressed on AWA neuron [43] because, together with sri-14, odr-10 also senses diacetyl (low concentrations versus high concentrations) and it is one of the few GPCRs linked to a specific target molecule [58, 59].

Unfortunately, since any experimental information about their structure is available, a bioinformatic prediction must be performed. We use the GPCR-I-TASSER [51] server protocol to do it and it returns the ten closest X-ray resolved structures to the predicted one. Almost every X-ray resolved structure has at least a “specific ligand” in its template, where by “specific ligand” we assume that this ligand has a pharmaco-biological significance (for example an inhibitor or an activator). We exclude templates with ligands with unclear significance or useful to our analysis (e.g. cofactors, metals, solvents etc). With these criteria, a collection of more than 20 resolved protein templates is thus retrieved. We clear each PDB file in order to have a clean “protein-ligand only” file, ready to be submitted to our algorithms.

Binding site definition

Given each GPCR-ligand complex the binding region was defined by calculating the distance between all C-alpha atoms of the protein and all the atoms of the small molecule. We consider a residue belonging to the real Binding Site (called BS) if it has at least one C-alpha atom with a distance less than 6 Å with at least one ligand atom.

For the purpose of comparison, we have adopted the following procedure for the definition of Binsing Site Decoy (called BSD):

- The aim is to define a set of randomly selected BSD on the receptor surface. To this end, we define the BSD in such a way that the number of residue that make up the BSD to be on average as close as possible to the number of BS residue.
- We center a “probe sphere” on each C-alpha of any receptor and we define for each residue a BSD which is dependent on the radius R of the sphere.
- For each residue Ri the BSD is composed of the residues that have a distance between their C-alpha and the C-alpha of the residue Ri less than R.
- The purpose of this procedure is to vary R in order to find the optimal value in terms of patch size. As R increases, the number of residues increases and the patch size tends to increase. The procedure starts with a radius of 6 Å and increasing its dimension by 1 Å each iteration step, until a maximum value of 20 Å.
- Given a GPCR, we define the BSDs considering the radius R which provides (on average) the number of residues belonging to the BSDs closest to the number of residues belonging to the BS. Therefore, each GPCR has the specific patch decoy size according to the size of the real binding region.

Surface patch analysis with Zernike descriptors

For each receptor and its cognate ligand, we calculate separately the molecular surface using DMS software [31]. We extracted the portion of protein surface regarding selected residues. Then, with a voxelization procedure, we represent the protein patch and the ligand as 3D functions.

These 3D functions can be expanded in the basis of the 3D Zernike Polynomials [30, 36, 44]: the norm of the expansion coefficients constitutes a ordered set of descriptors that compactly characterize the geometrical shape of the molecular surface.

Indeed, a function \( f(r, \theta, \phi) \) can be represented as:

\[
f(r, \theta, \phi) = \sum_{n=0}^{\infty} \sum_{l=0}^{n} \sum_{m=-l}^{l} C_{nlm} Z_{nlm}^m(r, \theta, \phi)
\]

where \( Z_{nlm}^m \) are the 3D Zernike polynomials, while the coefficients \( C_{nlm} \), are defined as Zernike moments.

Zernike polynomials can be written as:

\[
Z_{nlm}^m(r, \theta, \phi) = R_{nl}(r) Y_{l}^{m}(\theta, \phi)
\]

where the \( Y \) functions are complex spherical harmonics. These functions depend on both the angular variable \( \theta \) and \( \phi \), while the dependence on the radius variable, \( r \), is enclosed in the function \( R \), which is given by the following expression:

\[
R_{nl}(r) = \sum_{k=0}^{l-n} N_{nlk} r^{l-k} \]

where \( N \) is a normalization factor.

Therefore, the 3D Zernike moments of molecular surface (described by a function \( f(r, \theta, \phi) \)) are defined as the coefficients of the expansion defined in eq 3:
\[ C_{nlm} = \int_{|r| \leq 1} f(r) Z_{nlm}^*(r, \theta, \phi) dr \]  

(6)

where \( Z \) is the polynomial complex conjugate.

Their rotation invariant norms constitute the Zernike descriptors:

\[ D_{nl} = \| C_{nlm} \| = \sqrt{\sum_{m=-l}^{l} (C_{nlm})^2}. \]  

(7)

The detail of the Zernike formalism can be modulated modifying the order of the expansion \( n \). In our implementation, we set the maximum order of expansion to 10, giving a total of 36 invariants for each surface studied.

Represented as an ordered set of 36 numbers, the shape complementarity between 2 surfaces can be easily evaluated [32, 33, 38]. Since the shape of 2 perfectly fitting surface is exactly the same, they share very similar descriptors. Indeed, using the manhattan distance \( D(X, Y) = \sum_i |X_i - Y_i| \), when 2 surfaces have a low distance between them, they are characterized by a similar shape and therefore they are suitable for binding.

For each GPCR-ligand complex, we normalized the complementarity with the Z-score, as following:

\[ Z_i = \frac{X_i - \mu(X)}{\sigma(X)} \]  

(8)

where the subscript \( i \) refers to the \( i \)-th patch considered, \( x_i \) is the Manhattan distance between its Zernike descriptors and the ones of the ligand, \( \mu \) and \( \sigma \) represent the mean and the standard deviation of all the patch-ligand distances, respectively.

**Molecular dynamics simulations**

All simulations were performed using Gromacs 2020.6 [60]. Parameters of the systems were built using Swiss-Param web server [61]. Molecules was placed in a cubic simulative box, with periodic boundary conditions, filled with TIP3P water molecules [62]. All the molecules were simulated for 100 ns, using an integration step of 1 fs and saving molecules conformations each ns.

**Acknowledgements**  The research leading to these results has been also supported by European Research Council Synergy grant ASTRA (n. 855923).

**Data and software availability**  All the GPCR-ligand structures can be freely downloaded from the Protein Data Bank (PDB) [https://www.rcsb.org/]. The protein surface was calculated using with standard parameters dms software [31]. The protein structures analysis and the voxelization were performed using in-house R scripts, relying mainly on bio3d [64] and geometry [65] packages. The 3D Zernike descriptors calculation was done using the python code described in [66]. Code is available upon request.

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