

Molecular sampling of the allosteric binding pocket of the TSH receptor provides discriminative pharmacophores for antagonist and agonists

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Abstract

The TSHR (thyrotropin receptor) is activated endogenously by the large hormone thyrotropin and activated pathologically by auto-antibodies. Both activate and bind at the extracellular domain. Recently, SMLs (small-molecule ligands) have been identified, which bind in an allosteric binding pocket within the transmembrane domain. Modelling driven site-directed mutagenesis of amino acids lining this pocket led to the delineation of activation and inactivation sensitive residues. Modified residues showing CAMs (constitutively activating mutations) indicate signalling-sensitive positions and mark potential trigger points for agonists. Silencing mutations lead to an impairment of basal activity and mark contact points for antagonists. Mapping these residues on to a structural model of TSHR indicates locations where an SML may switch the receptor to an inactive or active conformation. In the present article, we report the effects of SMLs on these signalling-sensitive amino acids at the TSHR. Surprisingly, the antagonistic effect of SML compound 52 was reversed to an agonistic effect, when tested at the CAM Y667A. Switching agonism to antagonism and the reverse by changing either SMLs or residues covering the binding pocket provides detailed knowledge about discriminative pharmacophores. It prepares the basis for rational optimization of new high-affinity antagonists to interfere with the pathogenic activation of the TSHR.

Introduction

GPCRs (G-protein-coupled receptors) are major pharmacological targets for therapeutic applications [1]. Together with FSHR (follicle-stimulating hormone receptor) and LHCGR (luteinizing hormone/choriogonadotropin receptor), TSHR (TSH [thyroid-stimulating hormone; thyrotropin] receptor) is a member of the GPHRs (glycoprotein hormone receptors) within family A of GPCRs. TSHR and TSH are involved in a variety of physiological functions in the human body. Stimulation of the receptor leads to increased production and secretion of thyroid hormones, triiodothyronine and thyroxine. The thyroid hormones are primarily responsible for regulation of metabolism such as cell growth and proliferation [2] and are involved in embryonic and early postnatal brain development [3]. Beside its function in the thyroid gland, TSHR is expressed in multiple extrathyroidal tissues (bone, brain, kidney, adipocytes and immune system cells) [4–6]. However, the physiological role in these tissues is poorly known until now.

TSHR is activated endogenously by the large hormone thyrotropin and activated pathologically by auto-antibodies.

Key words: compound 52, constitutively activating mutation, G-coupled-protein receptor, small molecular ligands, thyrotropin receptor.

Abbreviations used: c52, compound 52; CAM, constitutively activating mutation; ECL, extracellular loop; GD, Graves' disease; GO, Graves' ophthalmopathy; GPCR, G-protein-coupled receptor; GPHR, glycoprotein hormone receptor; ICL, intracellular loop; LHCGR, luteinizing hormone/choriogonadotropin receptor; SML, small-molecule ligand; TMH, transmembrane helix; TSH, thyroid-stimulating hormone; TSHR, thyrotropin receptor; wt, wild-type.

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Such TSHR-stimulating antibodies induce constitutive activation of the receptor and cause overproduction of thyroid hormones leading to hyperthyroidism and GD (Graves' disease) [7]. One of the prominent symptoms is GO (Graves' ophthalmopathy). Current treatment options for Graves' hyperthyroidism and GO are inadequate because they are often invasive and generally target the signs and symptoms of the disease rather than the pathophysiology [8]. A potential approach could be based on the suppression of pathogenic autoantibody activation directly on the TSHR level by interference by drug-like SMLs (small-molecule ligands). In contrast with the activating antibodies and thyrotropin that bind to the TSHR extracellular region, synthetic SMLs bind within the transmembrane domain into an additional allosteric binding pocket. Recently, SMLs have been identified which bind in this allosteric binding pocket (reviewed in [9]). The design and application of a HTS (high-throughput screening) assay, especially designed for GPCRs to allow for quick and cost-effective testing of large compound libraries has been developed to identify such SMLs as novel drug candidates. However, the counterpart residues of these SMLs on their targets are studied only incompletely. Using site-directed mutagenesis the amino acids that are covering this allosteric binding pocket of TSHR were recently characterized [10,11].

First, the present review focuses on the recent discovery and characterization of two different mutant types (CAMs [constitutively activating mutations] and inactivating mutations) of residues covering the allosteric binding region. These

signalling-sensitive positions may be potential interaction points of allosteric ligands that may lead to inactivation or activation of the TSHR. Secondly, we examine SMLs that bind into this pocket, located within the transmembrane helical bundle.

Analyses of structure–function relationships between chemical structures of SMLs and functional effects caused by mutation are helpful to gain insights into the interaction between SMLs and wt (wild-type) amino acids covering the allosteric binding site of the TSHR. The knowledge about discriminative pharmacophores on the counterparts, on ligand as well as receptor site, facilitates the extraction of specific fingerprints for agonistic or antagonistic features of SMLs for the TSHR. The long-term aim of these studies is to develop high-affinity and selective antagonists that might have the potential to interfere with pathogenic activation of the TSHR as occurs in GD and GO.

Allosteric binding pocket in GPHRs

Family A GPCRs have been evolving for 570–700 million years (reviewed in [12]) and possess the general structural topology of an extracellular N-terminal region: seven TMHs (transmembrane helices) connected by three ICLs (intracellular loops) and three ECLs (extracellular loops) and a C-terminal tail. They bind diverse ligands such as amines, purines, lipids or peptides between the transmembrane helical bundle. This transmembrane binding pocket region of family A GPCRs is generally known to be sensitive to endogenous ligand binding [13–16], with the exception of GPHRs. The endogenous glycoprotein hormones of GPHRs bind to a large extracellular orthosteric binding site leading to receptor activation by triggering the transmembrane domain [17,18]. This orthosteric binding pocket is not interesting for pharmacology; on the one hand, the bulky hormone cannot be mimicked by drugs, on the other hand, their large extracellular orthosteric site turned out to be undruggable itself, owing to the large and rather flat binding area (reviewed in [19]). Truncations of the large N-terminal domain [20], mutagenesis studies in the transmembrane domain of TSHR [21] and binding studies of SMLs on LHCGR [22] provided evidence that drugs such as SML bind alternatively into an allosteric site located in the transmembrane helical bundle of GPHRs. Thus, although the orthosteric hormone-binding pocket in the TSHR and other GPHRs is located extracellularly, it appears that the binding pocket in the transmembrane region is also retained in these receptors at a comparable location as known for other GPCRs [23].

In previous studies, amino acid residues covering the TSHR-binding pocket have been characterized. Modelling-driven site-directed mutagenesis of amino acids lining this pocket led to the identification of two types of mutations: CAMs and inactivating mutations. CAMs are located in several helices: V421I (TMH1), Y466A (TMH2), T501A (TMH3), L587V (TMH6), M637C (TMH6), M637W (TMH6), S641A (TMH6), Y643F (TMH6), L645V (TMH6) and Y667A (TMH7). The wt amino acid residues showing

CAMs are expected to be key players in stabilizing the basal receptor conformation. These mutations indicate signalling sensitive positions and mark potential trigger points for receptor activation by agonists [11].

Of note, the mutation M637W at position 6.48 (Ballesteros and Weinstein numbering) in TMH6 has a significant effect on the interaction of the receptor with the SML agonist. At this position the strongest constitutive activation within the TMHs of TSHR was found. In the majority of family A GPCRs, tryptophan is located at position 6.48 and has been known to be involved in receptor activation. According to these data one can speculate that Met⁶³⁷ is involved in both stabilizing the basal and supporting the active conformation [11]. This indicates that this position is a potential interaction partner for agonists. However, this also implies that antagonists should avoid this activation-sensitive Met⁶³⁷ position.

The identified inactivating mutations, also termed silencing mutations, led to an impairment of basal TSHR activity. The TSHR three-dimensional homology model visualizes the spatial distribution of mutants within the binding pocket. Two spatial clusters of silencing mutations flanking the allosteric binding pocket have been described in [10]. Cluster I is arranged between amino acids Val⁵⁰² (TMH3), Leu⁵⁵² (TMH4), Tyr⁵⁸² (TMH5) and Met⁵⁷² at ECL2. Silencing mutations in the ECL2 were previously reported [24]. Amino acids Val⁴²⁴ (TMH1), Leu⁴⁶⁷ (TMH2) and Leu⁶⁶⁵ (TMH7) form cluster II. Such silencing mutations switch the receptor to a more inactive conformation. Potential antagonistic SMLs should be able to constrain the TSHR in an inactive conformation by interacting with these positions known for their ability to reduce the basal TSHR activity. Taken together, all these signalling-sensitive amino acids residues (extracted from <http://www.SSFA-GPHR.de> [25] and summarized in Table 1) indicate locations where SMLs may shift the receptor to either an inactive or an active conformation. CAMs mark contact points for agonists and silencing mutations mark the contact points for antagonists.

Allosteric modulators

The first small molecule interacting with the TSHR was originally developed as an agonist for the LHCGR [26]. Molecular modelling and functional experiments guided the chemical derivation of the LHCGR agonist and led to the identification of the first TSHR-specific partial agonist Org-41841 [27] as well as low-affinity antagonist c52 (compound 52) [20]. Both compounds share the same basic thienopyrimidine scaffold structure. This knowledge, from *in silico* as well as functional studies, directed the following investigations that resulted in the identification of several different TSHR ligands [23,28]. To gain a better understanding of the interactions of SMLs within the binding pocket we report about the effects of SMLs on signalling-sensitive amino acids at TSHR and use c52 and Org-41841 as examples. The two SMLs are very similar and differ only by an extended side chain on the aromatic ring of the antagonist (see Figure 1 and [27]). Based on our three-dimensional homology

Table 1 | Residues of the TSH receptor covering the allosteric binding pocket that show signalling-sensitive effects upon mutation either as constitutive activating mutation (CAMs) or as silencing mutation

Mutations known to cause disease-relevant effects are indicated for *inactivating pathogenic and for †activating pathogenic mutations.

Location	Ballesteros and Weinstein numbering	Amino acid human TSHR	Silencing mutation	CAM	Reference
TMH1	1.39	Val ⁴²¹		Ile	[11]
	1.42	Val ⁴²⁴	Ile		[10]
TMH2	2.53	Met ⁴⁶³		Val†	[29]
	2.56	Tyr ⁴⁶⁶		Ala	[11]
	2.57	Leu ⁴⁶⁷	Pro*,Val		[10,30]
	2.64	Asp ⁴⁷⁴	Glu		[31]
TMH3	3.32	Thr ⁵⁰¹		Ile	[32]
	3.36	Ser ⁵⁰⁵		Asn†, Arg	[33–37]
	3.38	Leu ⁵⁰⁷	Ser		[36]
	3.40	Val ⁵⁰⁹		Ala	[36]
ECL2		Ile ⁵⁶⁸	Leu	Ala, Phe, Val, Thr†	[24,38]
		Leu ⁵⁷⁰		Ala, Phe	[21,24]
		Met ⁵⁷²	Ala		[24,39]
TMH5	5.39	Tyr ⁵⁸²	Ala, Phe		[10]
	5.43	Val ⁵⁸⁶	Ile		[10]
	5.44	Leu ⁵⁸⁷		Val	[11]
TMH6	5.51	Phe ⁵⁹⁴	Ile		[10]
	6.45	Phe ⁶³⁴	Ile		[10]
	6.48	Met ⁶³⁷		Cys, Trp	[11]
	6.50	Pro ⁶³⁹		Ser†	[40,41]
	6.51	Ile ⁶⁴⁰	Leu	Lys†, Val	[24,42]
	6.52	Ser ⁶⁴¹		Ala	[11]
	6.53	Phe ⁶⁴²	Ile		[10]
	6.54	Tyr ⁶⁴³	Ala	Phe	[10,11]
TMH7	6.56	Leu ⁶⁴⁵		Val	[11]
	7.40	Leu ⁶⁶⁵	Val		[10]
	7.42	Tyr ⁶⁶⁷		Ala	[11]

model of the TSHR, the localization of both compounds within the binding pocket in the transmembrane domain was predicted and shown to be very similar, but differs in a shifted orientation of the molecules.

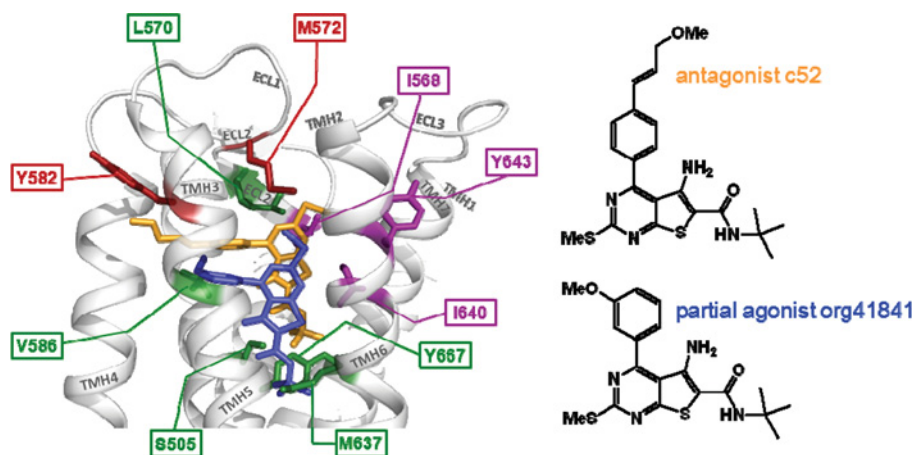
The partial agonist Org-41841 is smaller and binds deeper into the binding region and is therefore able to interact with the Met⁶³⁷ at position 6.48 on TMH6. This is consistent with the finding that position Met⁶³⁷ has been identified as a key player of activation where the strongest CAM was found [11]. In contrast, c52 revealed a slightly shifted localization towards the extracellular side in the binding pocket in comparison to the partial agonist Org-41841. Thus c52 cannot reach Met⁶³⁷. The shifted orientation of c52 within the binding pocket is the consequence of an enlarged substituent on the phenyl group of this compound. Moreover, this prolonged side chain extends into the region where more inactivating mutations were found. Docking studies predicted a sensitive amino acids residue Tyr⁶⁶⁷ at TMH7 (position 7.42) as a contact point for c52. In the molecular model, c52 is constrained by the Tyr⁶⁶⁷ residue, therefore c52 is not able to reach Met⁶³⁷.

Effect of allosteric modulators

Surprisingly, the antagonistic effect of SML c52 was reversed to an agonistic effect, when tested at the constitutively activating TSHR mutant Y667A. The side chain reduction by an alanine mutation releases c52 that then subsequently hits Met⁶³⁷, which is located at the bottom of the binding pocket at position 6.48, where a highly conserved tryptophan at TMH6 exists on other GPCRs. Met⁶³⁷ was identified as a key residue for activation by showing a very strong constitutive activation that indicates its potential as an interaction partner for agonists. These reverse effects confirm not only the predicted binding site for c52, but also provide details about distinguishing locations for antagonists and agonist. Docking studies of partial agonist Org-41841 into the three-dimensional TSHR homology model predicted a hydrogen bond interaction between the amino group of the ligand and residue Glu⁵⁰⁶ (TMH3). Interestingly, mutant E506A did not lead to receptor activation by stimulation with Org-41841, whereas the activation by TSH is not affected. The alanine substitution at this highly conserved position

Figure 1 | Homology model of TSHR

Allosteric binding pocket in between the TMHs. The bound partial agonist (blue) is interacting with residues that showed constitutive activation upon mutation (green residues). The low-affinity antagonist (orange) is docked with a shifted position more to the extracellular side. Agonist and antagonist are similar, differing only by the extended substituent on the aromatic ring. This substitution is pointing to residues where inactivating mutations were found upon mutation (red residues). Positions where both effects depend on the side chain exchange have been described and are highlighted in magenta.



Glu⁵⁰⁶ resulted in a breakdown of receptor activation. These data indicates ligand–receptor interaction and supports the predicted orientation of the ligand within the pocket [21].

In conclusion, elucidation of detailed activating and inactivating patterns among residues covering the allosteric binding site of TSHR allows for directed manipulations to switch from agonism to antagonism and vice versa. Modifications either at signalling-sensitive receptor positions at the allosteric binding pocket or at an SML can modify or even reverse the activation state of TSHR. This strategy revealed detailed knowledge about discriminative pharmacophores for agonistic and antagonistic features of SMLs, which can now be used to guide future studies. The information gained about complementary properties of the allosteric binding region and the small molecule ligands are the basis for the optimization of high-affinity antagonists for the TSHR for treatment of GO or constitutive activation by pathogenic mutations.

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