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Identification of Specific Transmembrane Residues and Ligand-Induced Interface Changes Involved In Homo-Dimer Formation of a Yeast G Protein-Coupled Receptor[†]

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ABSTRACT: The Saccharomyces cerevisiae α-factor pheromone receptor, Ste2p, has been studied as a model for G protein-coupled receptor (GPCR) structure and function. Dimerization has been demonstrated for many GPCRs, although the role(s) of dimerization in receptor function is disputed. Transmembrane domains one (TM1) and four (TM4) of Ste2p were shown previously to play a role in dimerization. In this study, single cysteine substitutions were introduced into a Cys-less Ste2p, and disulfide-mediated dimerization was assessed. Six residues in TM1 (L64 to M69) that had not been previously investigated and 19 residues in TM7 (T278 to A296) of which 15 were not previously investigated were mutated to create 25 single Cyscontaining Ste2p molecules. Ste2p mutants V68C in TM1 and nine mutants in TM7 (cysteine substituted into residues 278, 285, 289, and 291 to 296) showed increased dimerization upon addition of an oxidizing agent in comparison to the background dimers formed by the Cys-less receptor. The formation of dimers was decreased for TM7 mutant receptors in the presence of α-factor indicating that ligand binding resulted in a conformational change that influenced dimerization. The effect of ligand on dimer formation suggests that dimers are formed in the resting state and the activated state of the receptor by different TM interactions.

G protein-coupled receptors (GPCRs)¹ are membrane proteins that form one of the largest and most diverse families of proteins in eukaryotes ranging from yeast to human. Though the primary sequences are different among the GPCRs, all GPCRs share common structural features: seven transmembrane helical domains (TMs) across the lipid bilayer, with the TMs connected by intracellular and extracellular loops, an extracellular N-terminus, and an intracellular C-terminus (1). GPCRs mediate responses to various stimuli such as hormones, odors, peptides, and neurotransmitters. Binding of ligand to a GPCR triggers receptor-specific signals through a heterotrimeric G protein. Since it has been reported that genetic variation of GPCRs often alters receptor functions such as ligand binding, G protein coupling, and receptor life cycle, GPCR mutation is considered a causative agent of many of human diseases (2). GPCRs have been the most successful molecular drug targets in clinical

Ste2p is the α -factor pheromone receptor in Saccharomyces cerevisiae and has been used as a model for the study of the molecular basis of GPCR function (4-6). Ste2p can be replaced in yeast cells with mammalian receptors with functionality conserved (7), and Ste2p can be expressed and trigger signal transduction upon ligand binding in HEK293 cells (8). Also, Ste2p can serve as an established model for fungal GPCRs. Recently, many more GPCRs in fungi have been identified and classified into six different categories based on sequence homology and ligand sensing [for reviews see ref 9]. Ste2p is the most well studied receptor among fungal GPCRs, some of which are suggested to be related to fungal pathogenesis [for reviews see ref 9].

Recently, evidence has been growing that many GPCRs form homo- and/or heterodimeric or oligomeric complexes [for reviews see refs (9-11)]. Oligomerization has been discovered by techniques such as cross-linking, bioluminescence resonance energy transfer, fluorescence resonance energy transfer, and immunoprecipitation (10). Dimerization is thought to be important for various aspects of GPCR function such as receptor biogenesis, formation of ligand-binding sites, signal transduction, and down-regulation (11, 12). However, the view that dimers are involved in the rhodopsin-like (Class 1A) receptor-activated signaling has been challenged (13-16).

It has been demonstrated that Ste2p is internalized as a dimer/ oligomer complex (17, 18), and oligomerization-defective mutants can bind α -factor but signaling is impaired (19). It has also been shown that the dominant/negative effect on wildtype signaling of a signaling-defective mutation in Ste2p (Ste2p-Y266C) can be partially reversed by mutations in the G⁵⁶XXXG⁶⁰ dimerization motif, indicating that signal transduction by oligomeric receptors requires an interaction between functional monomers (20). Recently, dimer interfaces were

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Abbreviations: Cu-P, copperII-1,10, phenanthroline; EL, extracellular loop; G protein, heterotrimeric GTP-binding protein; GPCR, G protein-coupled receptor; FT-HT, Cys-less Ste2p with the FLAG and His epitope tags; FT-HT-Xa, the template construct or Cys-less Ste2p with FLAG Tag, His tag, and Factor Xa cleavage site; MLT, medium lacking tryptophan; MLU, medium lacking uracil; SDS-PAGE, sodium dodecyl sulfate polyacrylamide gel electrophoresis; Ste2p-FT-HT, Ste2p with FLAG and His epitope tags; SD medium, Synthetic defined medium; TM, transmembrane domain.

Table 1: Receptors Used in This Study						
receptor name	description					
wild-type Ste2p-FT-HT FT-HT FT-HT-Xa	native Ste2p wild-type with FLAG/His epitope tags Cys-less Ste2p-FT-HT FT-HT with tandem Factor Xa cleavage sites,					

template for all Cys mutants generated in this study

identified in Ste2p near the extracellular end of TM1 and TM4 (21). In that study it was found that dimerization was symmetric, occurring between receptors at the TM1-TM1 interface or the TM4-TM4 interface. In our current study, using the disulfide cross-linking methodology, we studied the participation of specific residues at the intracellular boundary between TM1 and intracellular loop one and the entire TM7 in Ste2p dimerization.

EXPERIMENTAL PROCEDURES

Strains, Media, and Plasmids. S. cerevisiae strain LM102 described by Sen and Marsh (22) was used in the growth arrest and LacZ assays. The genotype for the LM102 strain is MATa, bar1, his4, leu2, trp1, met1, ura3, FUS1-lacZ::URA3, ste2-dl (deleted for the α -factor receptor). The protease-deficient strain BJS21 (MATa, prc1-407 prb1-1122 pep4-3 leu2 trp1 ura3-52 ste2::Kan^R) was used in disulfide cross-linking and Western blot assays to decrease receptor degradation during analyses (23). The parental plasmid, pHY4 expressing the template construct used for mutagenesis, FT-HT-Xa (cys-less Ste2p with the FLAG and His epitope tags with Factor Xa cleavage cite, see Table 1 for description of the various receptor constructs used in this study) was generated by introducing a tandem Factor Xa cleavage site between Val192 and Thr193 into pBec2 expressing FT-HT (FLAG and His tagged Cys-less Ste2p) under a consititutive GPD promoter (24). Twenty-five single Cys mutations ranging from Leu⁶⁴ through Met⁶⁹ on TM1 and Thr²⁷⁸ through Ala²⁹⁶ on TM7 were generated in the pHY4 background by PCR-based site-directed mutagenesis (25). For coexpression experiments, plasmid pHY6 was constructed from p426GPD, a 2- μ m based shuttle vector with a GPD promoter, CYC1 terminator, and URA marker for selection in yeast (26). STE2 containing Cterminal FLAG and His epitope tags and a tandem Factor Xa digestion site in EL2 was PCR-amplified from plasmid pHY4 using primers that introduced BamHI and EcoRI restriction sites. The resulting PCR product was subcloned into the complementary sites of p426GPD. Also, to create pHY6 an epitope tag comprising codons encoding a nine-amino acid sequence of rhodopsin (rho-tag) was substituted for the FLAG and His epitope tags by ligation of a C-terminus part of STE2 product amplified from pBKY1(27). Cys mutants in this template were generated as described above. The sequences of constructs were verified by DNA sequence analysis completed by the Molecular Biology Resource Facility located on the campus of the University of Tennessee. Primers were purchased from Sigma Genosys or IDT (Coralville, IA). After sequence confirmation, constructs were transformed into the ste2-deletion strains LM102 and BJS21 by the method of Gietz et al. (28). Transformants bearing the pHY4 or pHY6 constructs were selected by their growth in the absence of tryptophan for pHY4 on MLT medium (medium lacking tryptophan) or in the absence of uracil for

pHY6 on MLU medium (medium lacking uracil) (29). All media components were obtained from Difco.

Growth Arrest (Halo) Assay. Growth arrest was measured as described previously (30). Briefly, filter disks were impregnated with 10-μL portions of peptide solutions at various concentrations and placed onto the overlay containing S. cerevisiae LM102 cells. The plates were incubated at 30 °C for 24–36 h and then observed for clear zones (halos) around the disks. The halo measured included the diameter of the disk. The normalized activity of each mutant was determined by comparing the halo size for the FT-HT-Xa receptor at 2 μ g of α -factor. All assays were carried out at least three times with no more than a 2-mm variation in halo size at a particular amount applied for each

FUS1-lacZ Gene Induction Assay. S. cerevisiae LM102 contains a FUS1-lacZ gene that is inducible by mating pheromone. Cells were grown overnight in SD (synthetic defined) medium (yeast nitrogen base medium (Difco) without amino acid) supplemented with the required amino acids at 30 °C to 5×10^6 cells/mL, washed by centrifugation, and grown for one doubling (hemocytometer count) at 30 °C. Induction was performed by adding 10⁻⁶ M of α-factor to 1 mL of concentrated cells (1 \times 10⁷ cells/mL). The mixtures were vortexed and, after incubation at 30 °C with shaking for 2 h, cells were harvested by centrifugation, and each pellet was resuspended and assayed for β -galactosidase activity (expressed as Miller units) in duplicate by using a β -galactosidase assay kit (Pierce) according to the manufacturer's instructions. The activity of each mutant was normalized by comparing β -galactosidase activity for the wildtype strain. The standard deviation was determined from three independent experiments.

Preparation of Membranes. Membrane preparation of Ste2p was carried out essentially as described previously (30). Cells were grown to log phase, and then 1×10^8 cells were harvested by centrifugation and lysed by agitation with glass beads in a lysis buffer containing 50 mM Hepes, pH 7.5, 1 mM EDTA, 10 μ g/mL phenylmethylsulfonyl fluoride, 2 μ g/mL leupeptin, and 2 μ g/mL pepstatin. The lysate was cleared by centrifugation at 2000g for 5 min, and then membranes were harvested by centrifugation at 15000g for 45 min. The membrane pellet was washed and then resuspended in 100 μ L of a buffer (pH 7.4) containing 10% glycerol, 50 mM Hepes, 0.15 mM NaCl, 2 mM CaCl₂, 5 mM KCl, 5 mM MgCl₂, 4 mM EDTA (30). The protein concentration was determined by the Lowry assay (Pierce), and the membrane preparation was stored at -20 °C overnight and used for further assay the next day.

Disulfide Cross-Linking with Cu-Phenanthroline. The 100 μ g of membrane protein preparation was treated with a fresh preparation (pH 7.4) of Cu(II)-1,10-phenanthroline (Cu-P; final concentration, 2.5 µM CuSO₄ and 7.5 µM phenanthroline). The reaction was carried out at room temperature for 30 min, terminated with 50 mM EDTA, and kept on ice for 20 min followed by adding Laemmli sample buffer. For study of the time course of cross-linking, Cu-P treated samples were aliquotted at 45 s, 2 min, 5 min, 10 min, 20 min, and 30 min before termination. In all cases where a high level of cross-linking was observed, the membranes were treated with Cu-P and the reaction was terminated by EDTA only and by EDTA and NEM to prevent cross-linking that might occur upon subsequent manipulation of the sample. No differences were found in the extent of crosslinking with the additional NEM treatment. In experiments designed to prevent disulfide bond formation, the membranes

Table 2: Biological Activities of Cys Mutant Receptors

		β -galactosidase activity		growth arrest activity c	binding assay ^d	
receptor FT-HT-Xa		basal activity ^a 1.0 ± 0.1	induced activity ^b 1.0 ± 0.1		B-max 1.0 ± 0.3	$K_{\rm d} ({\rm nM})$ 1.0 ± 0.3
T65C	0.9 ± 0.1	0.9 ± 0.1	1.08	1.2 ± 0.2	1.5 ± 0.7	
L66C	1.0 ± 0.1	1.3 ± 0.5	1.00	0.5 ± 0.1	0.5 ± 0.2	
	I67C	1.0 ± 0.1	1.1 ± 0.3	1.04	1.5 ± 0.1	2.2 ± 0.5
	V68C	1.0 ± 0.1	0.9 ± 0.3	1.00	0.9 ± 0.2	1.1 ± 0.4
	M69C	1.1 ± 0.1	0.7 ± 0.2	1.00	0.7 ± 0.1	1.0 ± 0.4
TM7	T278C	0.7 ± 0.1	0.9 ± 0.2	0.88	0.7 ± 0.1	0.7 ± 0.3
	T279C	1.1 ± 0.2	0.9 ± 0.1	1.08	0.4 ± 0.1	0.4 ± 0.1
	V280C	0.8 ± 0.1	1.3 ± 0.5	1.00	0.3 ± 0.1	0.5 ± 0.1
	A281C	0.5 ± 0.1	1.1 ± 0.3	n.d	0.4 ± 0.1	0.8 ± 0.2
	T282C	1.8 ± 0.5	0.9 ± 0.3	1.00	0.4 ± 0.1	0.9 ± 0.2
	L283C	1.6 ± 0.3	0.7 ± 0.2	0.96	1.7 ± 0.7	4.7 ± 1.3
	L284C	1.0 ± 0.1	0.8 ± 0.2	0.92	0.6 ± 0.1	1.1 ± 0.3
	A285C	1.4 ± 0.1	1.0 ± 0.1	0.88	0.7 ± 0.2	0.9 ± 0.2
	V286C	1.3 ± 0.1	1.0 ± 0.3	1.12	0.7 ± 0.1	1.9 ± 0.7
	L287C	1.7 ± 0.1	1.0 ± 0.3	1.01	0.8 ± 0.1	1.1 ± 0.1
	S288C	1.1 ± 0.1	1.0 ± 0.3	1.00	0.7 ± 0.1	4.1 ± 1.5
	L289C	1.4 ± 0.1	0.8 ± 0.2	0.92	1.3 ± 0.2	2.1 ± 0.6
	P290C	1.4 ± 0.1	1.0 ± 0.1	0.88	n.d.	n.d.
	L291C	1.3 ± 0.1	1.0 ± 0.3	0.88	0.6 ± 0.1	1.7 ± 0.8
	S292C	1.4 ± 0.1	1.0 ± 0.3	1.04	0.6 ± 0.1	0.2 ± 0.1
	S293C	1.5 ± 0.1	1.0 ± 0.3	1.00	1.2 ± 0.1	2.5 ± 0.4
	M294C	1.2 ± 0.1	1.2 ± 0.4	0.94	0.8 ± 0.1	0.9 ± 0.2
	W295C	1.7 ± 0.3	1.5 ± 0.3	1.01	1.3 ± 0.1	1.3 ± 0.1
	A296C	1.7 ± 0.4	2.5 ± 0.8	1.00	0.7 ± 0.1	0.4 ± 1.5

^aRelative activity (±standard deviation) compared with basal activity of FT-HT-Xa (Miller unit of FT-HT-Xa was 3.9). ^bRelative activity (±standard deviation) compared with induced activity of FT-HT-Xa at 1 μM of α-factor (Miller unit of FT-HT-Xa was 31.7). ^cThe relative halo size (growth arrest) was compared to the halo size of the FT-HT-Xa receptor at 2 μg of α-factor applied to a disk (the halo size of FT-HT-Xa was 24 mm). The standard deviation of the halo activity for all receptors was within ± 0.2 (three replicates). n.d. = not detected. ^dThe Bmax and K_d values are presented relative to those of the FT-HT-Xa receptor. The Bmax and the K_d were determined by saturation binding of radioactive α-factor according to the protocol in Experimental Procedures. n.d. = not detected, as the binding was not greater than background (B-Max of FT-HT-Xa was 3491 and K_d was 10.7 nM).

were treated with 5 mM of NEM for 20 min prior to incubation with Cu-P reagent. Alpha-factor or antagonist ([desW1,desH2]- α -factor) (10 μ M final concentration) were added to the membrane preparation and incubation was allowed to proceed for 30 min prior to Cu-P treatment in experiments performed to examine the influence of ligand on dimerization.

Factor Xa Digestion. The membrane protein preparation (40 μ g) was incubated with 0.4 unit of Factor Xa (Novagen) in Factor Xa cleavage buffer (0.1 M NaCl, 50 mM Tris-HCl, 5 mM CaCl₂, pH 8.0) containing 0.1% Triton X-100 for 30 min. Each sample was divided into two aliquots. The reactions were terminated by adding one-third the volume of Laemmli sample buffer (30% glycerol, 3% SDS, 0.01% bromphenol blue, 0.1875 M Tris, pH 6.8). To one aliquot β-mercaptoethanol (final 1%, v/v) was added for reducing conditions. Samples were analyzed by SDS-PAGE and Western blotting.

Western Blot. Immunoblot analysis of Ste2p was carried out as described previously (30). Each sample was incubated at room temperature and then separated on NuPAGE 10% Bis-Tris SDS—polyacrylamide gel (Invitrogen) using either nonreducing or reducing conditions and electrophoretically transferred to Immobilon-P membrane (Millipore Corp., Bedford, MA). The blot was probed with anti-FLAG M2 antibody (Eastman Kodak Co.) or 1D4 antibody (a monoclonal antibody for rho-tag, purchased from Flintbox, BC, Canada), and the bands were visualized with the West Pico chemiluminescent detection system (Pierce). Blots were imaged, and the total density of all Ste2p

bands in each lane was determined using a ChemiDoc XRS photodocumentation system with Quantity One one-dimensional analysis software (Bio-Rad). The intensity of the monomer and dimer signals was measured by densitometry, and the percentage of dimer was calculated as [dimer/(dimer + monomer) \times 100]. The averages of the ratio were measured from at least three independent experiments and standard deviations are presented in Table 3.

Saturation Binding Assay with $[^3H]\alpha$ -Factor. Tritiated α -factor (10.2 Ci/mmol) (31) was used in saturation binding assays on total membrane preparations as described previously (29). Specific binding data were analyzed by nonlinear regression analysis for single-site binding using Prism software (GraphPad Software, San Diego, CA) to determine the B_{max} value (receptors/cell) for each mutant receptor. Each experiment was carried out at least three times. The close similarity among three replicates is indicated by the standard deviations shown in Table 2.

HIS-Select HC Nickel Affinity. Membrane preparations (500 μ g of protein) from each sample were resuspended in a solubilization buffer (50 mM Tris HCl, 150 mM NaCl, 1% Triton X-100, 5 mM imidazole, pH 8.0) overnight. The solubilized proteins were then mixed with HIS-select HC nickel affinity gel (Sigma) and incubated at 4 °C for 30 min with end-over-end mixing. The nickel gel was separated from the mixture by centrifugation and then washed five times with buffer (50 mM sodium phosphate, pH 8.0, 0.3 M NaCl, 5 mM imidazole). Ste2p

was eluted from the nickel gel two times using elution buffer (50 mM sodium phosphate, pH 8.0, 0.3 M NaCl, 250 mM imidazole). The eluted samples were analyzed by SDS-PAGE silver staining and Western blotting using antibodies against FLAG, a

Table 3: Effect of Ligand Binding on Cu-P(Cu(II)-1,10-phenanthroline) Stimulated Disulfide Bond Formation^a

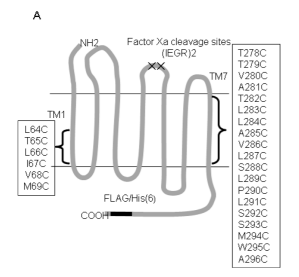
receptor		no treatment	Cu-P ^b	α, Cu-P ^c	A, Cu-P ^d
FT-HT-Xa		15.0 ± 4.4	12.0 ± 0.1	13.1 ± 0.2	12.2 ± 0.1
TM1	L64C	15.8 ± 0.1	11.3 ± 0.4	14.6 ± 1.0	13.6 ± 0.8
	T65C	16.2 ± 0.1	14.4 ± 0.1	15.9 ± 0.3	12.8 ± 0.3
	L66C	12.3 ± 0.4	11.9 ± 1.0	13.2 ± 0.4	11.9 ± 0.8
	I67C	19.3 ± 4.5	17.0 ± 0.1	17.4 ± 4.1	18.0 ± 2.0
	V68C	28.5 ± 3.2	99.2 ± 0.1	98.3 ± 0.8	98.8 ± 0.1
	M69C	17.7 ± 6.9	14.2 ± 5.0	17.0 ± 6.4	28.9 ± 10.2
TM7	T278C	15.1 ± 0.8	65.8 ± 8.8	12.5 ± 0.1	12.4 ± 0.1
	T279C	17.3 ± 0.3	10.1 ± 2.3	15.8 ± 0.2	10.9 ± 2.7
	V280C	15.3 ± 2.1	14.7 ± 1.5	16.3 ± 0.6	12.4 ± 0.1
	A281C	16.8 ± 0.5	11.9 ± 0.6	14.3 ± 1.5	11.3 ± 0.1
	T282C	11.7 ± 0.3	15.4 ± 0.1	12.0 ± 0.8	11.7 ± 0.1
	L283C	16.6 ± 0.4	15.7 ± 0.2	13.3 ± 0.1	10.7 ± 3.5
	L284C	15.2 ± 0.1	12.0 ± 0.2	17.7 ± 3.0	15.3 ± 0.1
	A285C	16.5 ± 0.8	53.5 ± 8.3	16.5 ± 2.1	17.7 ± 0.5
	V286C	15.6 ± 0.4	17.0 ± 0.3	13.2 ± 2.0	15.9 ± 0.4
	L287C	16.0 ± 0.1	14.3 ± 1.3	11.3 ± 1.7	16.0 ± 1.8
	S288C	14.7 ± 1.0	14.7 ± 1.5	11.5 ± 0.6	12.6 ± 0.2
	L289C	26.8 ± 2.0	61.0 ± 9.8	50.2 ± 12.9	40.9 ± 8.2
	L291C	31.8 ± 0.4	66.4 ± 0.1	52.0 ± 8.8	44.5 ± 5.6
	S292C	30.2 ± 8.3	82.3 ± 0.4	75.5 ± 11.4	65.9 ± 0.2
	S293C	21.1 ± 0.7	70.2 ± 5.3	38.5 ± 8.3	44.0 ± 0.1
	M294C	27.0 ± 7.6	75.6 ± 1.0	57.8 ± 0.7	50.8 ± 1.0
	W295C	15.7 ± 0.2	95.2 ± 0.7	76.2 ± 0.6	86.3 ± 0.1
	A296C	14.8 ± 2.1	91.1 ± 1.0	73.6 ± 1.5	82.1 ± 0.8

 $^a\mathrm{The}$ intensity of the monomer and dimer signal was measured by densitometry and the ratio of the signals was determined. The percentage of dimer was calculated as [dimer/(dimer + monomer) \times 100]. $^b\mathrm{Cu-P} = \mathrm{Cu-P}(\mathrm{Cu}(\mathrm{II})\text{-}1,10\text{-phenanthroline})$ treatment. $^c\alpha$, Cu-P = membranes treated with alpha-factor and Cu-P(Cu(II)-1,10-phenanthroline). $^d\mathrm{A}$,Cu-P = membranes treated with antagonist and Cu-P(Cu(II)-1,10-phenanthroline).

nine-amino acid sequence of rhodopsin (rho-tag), or an antibody to the N-terminus of Ste2p (32).

RESULTS

Expression and Biological Activities of Cys Mutant Receptors. To begin the analysis of Ste2p dimerization, we chose 12 residues, 6 in TM1 and 6 in TM7 (L289-M294) proximal to the cytoplasmic face of Ste2p (Figure 1A) for mutation to Cys. To extend the initial cross-linking results, 13 additional residues in TM7 were mutated (T278C-S288C, W295C, and A296C) (Figure 1A). These targeted residues were chosen for several reasons: (i) The TM1 region chosen was one helix turn away from the sequence G⁵⁶XXXG⁶⁰ previously established as being involved in Ste2p dimerization (19), (ii) TM7 contains the AXXXA motif also suggested to play a role in dimerization of other proteins (33) and 20 known growth factor receptors with tyrosine kinase activity (34), (iii) eight of the TM7 residues mutated (S288-S293,W295 and A296) are conserved among fungal pheromone receptors (35), and (iv) TM1 and TM7 are in close proximity to one another in crystal structures of GPCRs (36, 37) and a model of Ste2p (35). To eliminate nonspecific cross-linking the template for these mutations was a Cys-less receptor. This template (FT-HT-Xa) also contained two C-terminal epitope tags (FLAG and 6XHis) and tandem Factor Xa cleavage sites (IEGRIEGR) in the second extracellular domain in order to facilitate detection of interdomain crosslinking (Figure 1A). Wild-type, FT-HT (Cys-less Ste2p with the FLAG and His epitope tags), and FT-HT-Xa (the template construct) receptors used in this study demonstrated almost identical biological activities in a growth arrest assay indicating that incorporating the protease site and the epitope tags did not alter receptor function (Figure 1B). In addition, FT-HT-Xa and Ste2p-FT-HT showed identical expression (data not shown). The expression level of each single-Cys mutant receptor was determined by Western blot analysis. All mutants except P290C showed several bands between 44 and 55 kDa and expression levels similar to that of FT-HT-Xa (Figures 2A and 4A). The



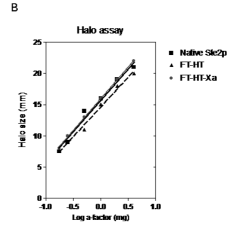


FIGURE 1: (A) The two-dimensional topology of *Saccharomyces cerevisiae* Ste2p. The cartoon indicates the location of the genetically engineered Factor Xa digestion site in EL2 and the epitope tags in the C-terminus. The two endogenous Cys residues (C59 and C252) were mutated to Ser to generate FT-HT-Xa which was the parental GPCR used to generate the TM1 and TM7 cysteine mutations in positions 64 to 69 in TM1 and 278 to 296 in TM7 as indicated in the boxes. (B) Dose—response analysis of growth arrest assay. The zone of growth inhibition of strains carrying the indicated receptors was measured at various concentrations of α -factor. FT-HT-Xa is the Cys-less receptor containing C-terminal FLAG and His epitope tags and a tandem Factor Xa digestion site in EL2. FT-HT is the Cys-less receptor without the Xa digestion site containing the FLAG and His epitopes, and Native Ste2p is the wild-type receptor that has no epitope tags and has two Cys residues (C59 and C252).

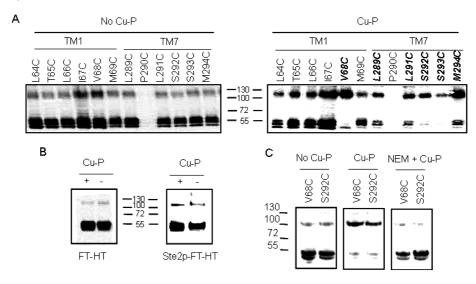


FIGURE 2: Effect of Cu-P (Cu(II)-1,10-phenanthroline) treatment on the dimerization of Ste2p containing single cysteine mutations. Membrane proteins were treated with Cu-P or left untreated, followed by SDS-PAGE, then immunoblotted and probed with anti-FLAG antibody. The upper band (~110 kDa) represents dimerized receptor and the lower band (~55 kDa) represents monomer. (A) Receptors with Cys in TM1 or TM7 were treated with Cu-P (right panel) or were untreated (left panel). Receptors indicated in bold exhibited a shift in signal from monomeric to dimeric form. (B) Controls with Cys-less Ste2p-FT-HT and Ste2p-FT-HT. (C) V68C and S292C receptors were untreated or treated with Cu-P in the presence or absence of NEM (N-ethylmaleimide) added prior to Cu-P treatment.

multiple bands are typical of Ste2p expression and are due to differences in glycosylation state, which does not influence receptor function (38). Although two intrinsic Cys residues have been substituted, a weak band at 110 kDa, corresponding to a dimerized form of Ste2p, was observed for the FT-HT receptor (Figure 2B). This band is likely a native, noncovalent dimer which was not disrupted by membrane protein preparation or SDS-PAGE. Such dimers have been observed on SDS-PAGE gels by other investigators working with Ste2p (20, 39, 40) and are also seen with the FT-HT-Xa receptor (Figure 5A; lane 1). This weak dimer observed in the Cys-less receptor is consistent with immunoprecipitation results showing that the two intrinsic Cys are not involved in dimerization (18). Membrane expression of P290C receptor was very low when judged by Western blot (data not shown). It is known that one of the roles of the conserved proline in TM7 in GPCRs is correct folding (41). The reduced expression of P290C was observed previously when it was shown that this receptor was primarily defective in plasma membrane localization using a P290C-GFP receptor, but the P290C-GFP construct did not demonstrate a defect in signaling (42).

Biological activity of each mutant was measured by growth arrest, β -galactosidase activity, and binding assays and is normalized relative to the parent receptor (Table 2). Most mutant receptors showed similar β -galactosidase activity and growth arrest activity as compared to FT-HT-Xa. Two mutants (M69C and L283C) showed slightly lower induced β -galactosidase activities and A281C did not show growth arrest activity, which is consistent with a previous study (43). The difference between β -galactosidase activity and cell division arrest has been observed in many mutagenic studies of Ste2p (44-46). It has been proposed that a short-term signaling effect is measured by the β -galactosidase assay, whereas the long-term effect is measured by the halo assay. The α -factor binding affinities of most Cvs mutants were within the experimental error range (± 2.5 -fold), with the exceptions of the S292C, receptor which showed an increase in binding affinity (5-fold lower K_d value), and the L283C and S288C which showed a decrease in binding affinity (up to 4.7-fold higher K_d values) as compared to that of FT-HT-Xa. B-Max values determined by a saturation binding assay indicated that the numbers of binding sites were similar to FT-HT-Xa for most mutant receptors, with the exceptions of T279C, V280C, A281C, and T282C, which showed expression about 50% lower than that of FT-HT-Xa. The P290C receptor showed no detectable alpha-factor binding, consistent with lack of expression in the Western blot analysis. Nonetheless, this receptor exhibited comparable growth arrest and β -galactosidase activities, suggesting that there were sufficient receptors expressed on the cell surface to elicit these biological responses upon ligand treatment (Table 2). Previous studies have shown that remarkably low levels of Ste2p at the cell surface are sufficient to manifest the biological responses (47). Taken together, we conclude that Cys mutation of the targeted residues did not severely interfere with receptor expression and function. We did not use the P290C receptor further in this study as it was expressed at a very low level.

Dimerization of Some Cys Mutants in TM1 and TM7 Is Markedly Enhanced by the Oxidizing Reagent Cu(II)-1,10-Phenanthroline. Initially, we tested the involvement of 12 TM residues, 6 on TM1 and 6 on TM7 (L289-M294) proximal to the cytoplasmic face of Ste2p (Figure 1A), in dimerization. As stated above, FT-HT-Xa and virtually all of the single Cys containing Ste2p mutants exhibited weak bands at about 110 kDa consistent with a small amount of dimerized Ste2p. However, in all cases the predominant band was near 55 kDa, the molecular mass of monomeric Ste2p. When these same receptors were treated with Cu(II)-1,10-phenanthroline (Cu-P), a reagent that has been used as an oxidizing reagent to drive disulfide bond formation in membrane proteins (48-51), many of the Cys mutated receptors showed marked increases in the 110 kDa band and a concomitant decrease in the monomer band (Figure 2A, Table 3). Notably, V68C, a TM1 mutant, and five mutants on TM7 (L289C, L291C, S292C, S293C, and M294C) showed strong (>60%) dimerization upon Cu-P treatment (Figure 2A, Table 3). When FT-HT (epitope tagged Cys-less receptor) and Ste2p-FT-HT (epitope tagged receptor with intrinsic cysteine residues at C59 in TM1 and C252 in TM5)

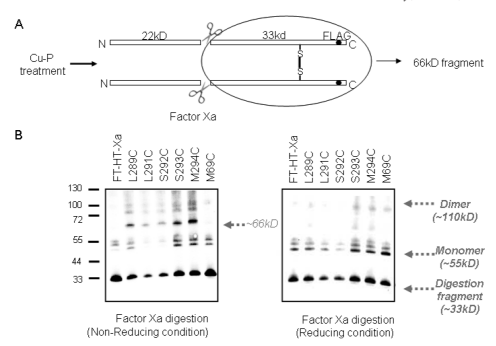


FIGURE 3: Protease Factor Xa digestions of single cysteine containing Ste2p receptors. (A) Diagram showing possible explanation for \sim 66 kDa band from TM7–TM7 interaction. (B) Total membrane proteins derived from cells expressing indicated receptors. Cu-P treated membranes with Ste2p containing single Cys mutations at various positions as indicated were treated with Factor Xa as described in Experimental Procedures. The digests were subjected to SDS–PAGE under reducing or nonreducing conditions, immunoblotted, and probed by anti-FLAG antibody.

receptors were tested, there was no difference in dimerization in the presence and absence of Cu-P treatment (Figure 2B) indicating that increased dimerization of mutant receptors was due to the specific cysteine residues engineered into Ste2p. To further test whether dimerization was mediated by disulfide bond formation involving cysteine residues, NEM (N-ethylmaleimide) pretreatment was used. NEM alkylates the free -SH group of cysteine irreversibly, so that disulfide bond formation cannot occur after NEM treatment. As shown in Figure 2C, pretreatment of V68C or S292C with NEM completely eliminated the Cu-P mediated dimerization. Similar results were found with the other TM7 residues tested (data not shown). In contrast to these six residues, placement of cysteine at positions 64, 65, 66, 67, and 69 followed by Cu-P treatment did not result in a dimer population that exceeded 20% (Figure 2A, right panel). Taken together, the results indicate that insertion of cysteine residues at position 68 on TM1 and positions 289, 291, 292, 293, or 294 on TM7 led to high levels of Ste2p dimerization through disulfide formation.

Although the studies described above provided evidence for Cu-P catalyzed Ste2p dimerization, it was possible that the single Cys mutant Ste2p is cross-linking with other proteins with similar molecular weight to yield the observed high molecular weight bands. To confirm that the observed dimer band in the Western blot was a Ste2p-Ste2p homodimer, we took advantage of the protease (Factor Xa) digestion site engineered into EL2 of Ste2p. Cross-linking between two Cys residues in TM7 followed by Factor Xa digestion would yield a homodimeric complex of 66 kDa (Figure 3A). Indeed, TM7 mutants (L289C, L291C, S292C, S293C, and M294C) digested with Factor Xa led to the appearance of a band near 66 kDa when SDS-PAGE was run under nonreducing conditions (Figure 3B; left panel). This band was markedly reduced when the gel was run in the presence of β mercaptoethanol, a potent disulfide reducing agent. The 66 kDa band was specific for dimerized receptors with a disulfide connection between monomers as the band was not observed

for FT-HT-Xa (Cys-less) and a TM1 mutant (M69C) in which there was no disulfide-mediated dimer formation (Figure 3B). The monomer bands (\sim 55 kDa) are due to an incomplete Factor Xa digestion. We performed partial digestion because a longer incubation led to the degradation of all bands. Bands at \sim 33 kD are Factor Xa digestion products of the monomeric receptor species (remaining in the samples). The generation of this band from the S292C and S293C mutant receptors, in which the majority of receptors formed dimers (Figure 2A, right panel), is likely the result of detergent-induced (Triton X-100 in Factor Xa digestion buffer) dissociation of background dimers. Indeed, the presence of background dimers from all the receptors tested including FT-HT-Xa (Cys-less) and M69C were reduced in the presence of Triton X-100. Further support for the disulfide bond induced dimer formation was provided when a Rho tagged M294C receptor coexpressed with a HIS tagged M294C receptor was retrieved as a dimer after pull-down using anti-HIS antibody both with and without Cu-P (data not shown). These results and the cross-linking results demonstrate the proximal location of the residues (L289-M294) on the intracellular parts of the TM7 domains of two Ste2p molecules and provide evidence for homodimerization of Ste2p involving TM7 as well as TM1.

The finding that six consecutive residues in TM7, with the exception of P290C, showed an increase in dimerization was unexpected since this region is believed to be α-helical. To see whether dimerization would occur throughout TM7, we expanded Cys replacement to include the full TM7 by generating 11 additional cysteine mutants N-terminal to the L289 residue (T278C, T279C, V280C, A281C, T282C, L283C, L284C, A285C, V286C, L287C, and S288C) and two cysteine mutants C-terminal to the M294 residue (W295C, A296C). Membranes from each mutant were prepared and processed with or without Cu-P treatment. Compared to the Cys-less FT-HT-Xa receptor, the T278C and A285C receptors exhibited at least a 3-fold increase in dimerization (Figure 4). Though dimer formation on the T278C mutant was not observed in a recent study (21), this mutant

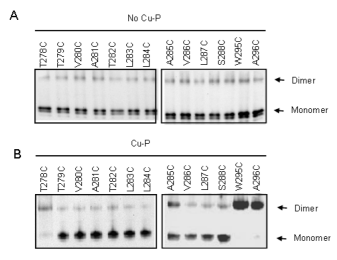


FIGURE 4: Effect of Cu-P (Cu(II)-1,10-phenanthroline) treatment on FT-HT-Xa containing cysteine replacements in TM7. Membranes were prepared, treated without (A) or with (B) Cu-P reagent, solubilized, and separated on SDS-PAGE. Each sample was immunoblotted and probed with anti-FLAG antibody.

showed strong dimer formation with Cu-P treatment in our study. The difference observed for this one residue is likely the result of variations in experimental design. In our current study Cu-P was used at a final concentration of 2.5 μ M, while in the previous study the concentration was $500 \mu M$. In addition, in our study we used membrane preparations, whereas whole cells were used by Wang et al. Receptors W295C and A296C showed a large increase in dimerization upon treatment with Cu-P compared to FT-HT-Xa (Figure 4A; Table 3). In contrast, nine mutants (T279C-L284C, V286C, L287C, and S288C) gave virtually identical dimerization results in the presence and absence of the oxidizing agent (Figure 4B; Table 3). These results indicate that unlike the residues C-terminal to Pro290, residues located N-terminal to Pro290 exhibit a periodicity with respect to dimer formation of Cys mutants which is consistent with the α-helical structure predicted for typical transmembrane domains of GPCRs. Because of a concern that disulfide cross-linking might trap and thereby favor the accumulation of dimer during the 30 min oxidization reaction, we analyzed the efficiency of the dimer formation at different times (Figure 5). The results indicated that L291C, S292C, S293C, M294C, and W295C started to form dimers after only 45 s of Cu-P incubation and reached halfmaximum levels within 5 min. Previously, the relative distance between TM residues in rhodopsin was estimated by the reaction time for disulfide formation between to Cys residues on different TMs (52). Thus, our data indicate that TM7 C-terminal to P290 is highly flexible allowing all of the residues in this part of TM7 to form disulfide bonds with more or less the same rapidity.

Effect of Ligand Addition on Dimerization. It is generally believed that activation of GPCRs upon ligand binding results in a conformational change involving rearrangement of transmembrane domains (53-56). To ascertain whether this conformational change would affect Ste2p dimerization, we investigated the changes in the dimerization pattern of Ste2p receptors in the presence and absence of ligand. The membranes expressing each mutant receptor were incubated with agonist (α -factor) or an antagonist (α -factor) and then treated with Cu-P. Dimer and monomer formation were monitored by Western blots (Figure 6A,B) and the percent dimer formation was determined (see Experimental Procedures, Table 3). Dimer formation of the Cys-less FT-HT-Xa receptor in the presence

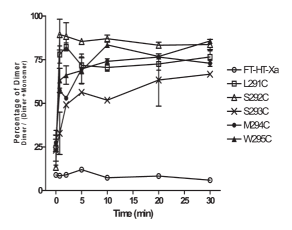


FIGURE 5: Time course analysis of dimer formation at residues C-terminal to P290 of TM7. Total membranes were prepared from cells expressing each receptors and treated with Cu-P (Cu(II)-1, 10-phenanthroline) for 45 s, 2 min, 5 min, 10 min, 20 min, and 30 min. The samples were analyzed by SDS-PAGE, followed by immunoblotting with anti-FLAG antibody. The average percentage of dimer population from three independent experiments was calculated and plotted.

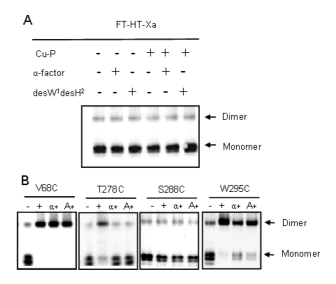


Figure 6: Effect of ligand binding on Cu-P (Cu(II)-1,10-phenanthroline) stimulated disulfide bond formation. (A, B) Total membrane protein was derived from cells expressing indicated receptors. Each sample was immunoblotted and probed with anti-FLAG antibody. FT-HT-Xa receptor was treated as indicated B "—" indicates no treatment, "+" indicates Cu-P treated samples, " α +" indicates samples incubated with α -factor followed by Cu-P treatment, and "A+" indicates samples incubated with the antagonist [desW1, desH2] α -factor followed by Cu-P treatment.

or absence of Cu-P was not affected by agonist or antagonist (Figure 6A and Table 3). Also, the ratio of dimer to monomer formed by Cu-P mediated cross-linking of the TM1 cysteine mutants did not change significantly after incubation with α -factor or an α -factor antagonist (representative results for V68C are shown in Figure 6B and Table 3). A time course of cross-linking at 0.75, 2, 5, 10, 20, and 30 min was also performed with the V68C receptor incubated with either α -factor or antagonist. The reaction was complete at 45 s (0.75 min), and no change in cross-linking relative to that observed in the absence of ligand was observed (data not shown). TM7 mutants which did not exhibit increased levels of dimerization upon Cu-P treatment (T279C-L284C, V286C, L287C and S288C), also showed no difference in dimer formation in the presence of α -factor or the

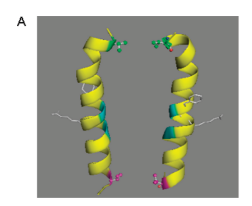
 $\alpha\text{-factor}$ antagonist (Table 3; raw data for S288C is shown in Figure 6B). However, for those TM7 receptors which exhibited a significant increase in dimer formation upon Cu-P treatment (T278C, A285C, L289C, L291C, S292C, S293C, M294C, W295C, and A296C), ligand treatment decreased the relative amount of dimer formation (Table 3; raw data for T278C and W295C are shown in Figure 6B). Taken together, the results suggest that the dimer interface of TM7 of Ste2p is changed in response to either agonist or antagonist binding.

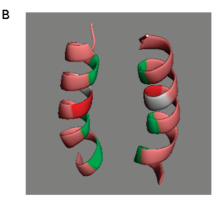
DISCUSSION

In this study, using a disulfide cross-linking methodology, we identified a specific residue in TM1 that interacts in Ste2p dimers, and we present the first evidence that residues in TM7 of this receptor participate in its dimerization. All of our studies were conducted with Ste2p in its membrane-bound state. Previously, it was shown that the maximum distance between α -carbons linked by disulfide bonds is about 7 Å (57). Thus, cross-linking experiments should identify amino acid side chains that are in close proximity when using the Cu-P oxidation reagent, which facilitates oxidation of sulfhydryl groups in cysteine residues. In analyzing our data, it is important to note that disulfide cross-linking might trap transient intermediates. Depending on the time of cross-linking relative to the rates of interconversion of the monomeric and dimeric states of the receptor, covalent

cross-linking might affect the equilibrium and bias the sample to yield more dimeric species than are present in the native population of the receptor. However, by comparing the cross-linked population of a receptor mutated in a specific region we believe we can learn about the relative tendencies of individual residues to participate in receptor-receptor contacts. Cross-linking between cysteine residues engineered into GPCRs and the use of Cu-P as an oxidizing reagent to facilitate disulfide bond formation between TMs in GPCRs has been used extensively by the Wess and Oprian laboratories (48, 51, 55, 58, 59) and was recently applied to Ste2p dimerization (21).

It had been suggested that TM1 formed a component of the interface between the two receptors in the Ste2p dimer (19, 21, 60). Our results support these findings and furthermore provide direct evidence for the involvement of V68 in TM1 in Ste2p dimerization. The mutant receptor V68C showed markedly increased dimerization over that of the FT-HT-Xa after Cu-P treatment, while under identical conditions all of the remaining TM1 Cys receptors showed at most a minor increase in dimer formation (Figure 2A; Table 3). The ability of V68C to form dimers is in good agreement with the recently published finding of a V45C-V45C cross-linking (21) since V45 and V68 are both located on the same face of TM1 and on opposite ends of TM1 (Figure 7A). It was previously proposed that G⁵⁶XXXG⁶⁰ residues in TM1 of one GPCR monomer interacted with a





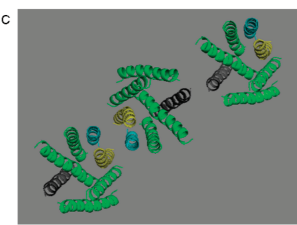


FIGURE 7: Dimerization interfaces of Ste2p. (A) V68 (red), a dimer contact found in the current study, is located on the same plane with V45 (green), a dimer contact at the extracellar end of TM1 (21). The backbones of two glycine residues (blue) known to play a role in Ste2p dimerization are shown. Two residues (F55, R58) involved in ligand binding are presented as a stick model located on the opposite side of the dimer interface. (B) TM7/TM7 dimer interface. The backbone of three dimer contacts identified from the current study (T278, A285, and L289, from top to bottom) are colored green. A281 (light gray) and T282 (red) would appear to be candidates for dimer contact based on a sideview of TM7 (Figure 8A), but the model here shows that alignments of these two residues are not favorable for disulfide bond formation. (C) Proposed oligomerization interfaces of Ste2p. Dimerization of Ste2p involving TM1/TM1 (magenta) and/or TM7/TM7 (yellow) interface allows TM4/TM4 interface (dark gray) to mediate a new dimerization contact enabling oligomerization. An extracellular view of a representative trimer is shown.

hydrophobic-rich surface of TM1 (perhaps involving residues 153, V57, A61, and L64), on the second monomer by a kind of "groove-in-ridge" association with Gly being the "groove" residues and the hydrophobic residues acting as the "ridges" (19). In this study, we found that V68C formed a disulfide bond. The fact that only one residue in the sequence (L⁶⁴T⁶⁶L⁶⁷I⁶⁷V⁶⁸M⁶⁹) formed this linkage suggests that the Ste2p-Ste2p interactions involving this region of the TM1 helix have significant spatial restrictions and that the TM1 helix at the carboxyl side of the G⁵⁶XXXG⁶⁰ sequence may be relatively rigid. We note that a biophysical analysis of TM1 in micelles indicates that the G⁵⁶XXXG⁶⁰ motif itself is flexible [ref 61 and unpublished results] and may facilitate TM-TM interactions between proximal helical elements. In any event, our mutational analysis of the TM1 domain defines a specific residue (V68) that appears to be involved in Ste2p dimerization.

In a previous FRET analysis, Overton and Blumer (60) analyzed the transmembrane domains of Ste2p that are involved in dimerization by expressing various combinations of Ste2p fragments. They concluded that TM1 is necessary and probably sufficient for dimerization, although they speculated that the N-terminus and TM2 may contribute to stabilize the dimers. However, our disulfide cross-linking data demonstrated that TM7 is also involved in the interface for dimerization of Ste2p. In the present study, we have analyzed full-length receptors, and it is possible that the fusion of fluorescent protein at the C-terminal of truncated receptors in the previous FRET studies hampered the interactions between TM7 domains. The involvement of TM7 in dimer formation is consistent with our previous observation that a TM6–TM7 fragment of Ste2p runs as a dimer in SDS–PAGE gel (62).

The finding that TM7 residues are involved in Ste2p dimerization leads us to propose that at least three dimerization interfaces can exist in Ste2p. In addition to the TM1 and TM4 interfaces previously found (21, 60), our data suggest that TM7-TM7 interactions are also involved in direct contacts in the Ste2p dimer. Since TM1-TM1 and TM4-TM4 contacts have been previously suggested to be involved in higher order oligomers (21) and a Ste2p trimer was demonstrated by cross-linking in a gel and by atomic force microscopy (40), it is possible that a single receptor can interact simultaneously with two additional receptors with different interfaces between the monomers: TM7 and/or TM1 can interact with TM7 and/or TM1 of a second receptor, while TM4 can interact with TM4 of a third receptor to form oligomers. A representation of a trimer is shown in Figure 7C with higher oligomers envisaged to contain additional interactions. On the basis of the existing experimental data it is not possible to conclude whether TM1-TM1 and TM7-TM7 dimers can be formed simultaneously in the cell.

The results described in this study show that cysteine residues introduced in positions T278, A285, and L289 on the extracellular half of TM7 form a disulfide bond with their counterpart in another Ste2p monomer (Figure 8A). This finding provides valuable information relating to the arrangement of the TM bundles of Ste2p and allow us to present a helical wheel projection of TM7 (Figure 8B) in which these three residues are oriented outward instead of facing inside the TM bundles as proposed previously in the Ste2p model based on rhodopsin (35). A281 and T282, which in a two-dimensional representation (Figure 8A,B) appear to lie on the same helix face, do not form dimers. However, our 3D model suggests that A281 and T282 are not close enough to form disulfide bonds when replaced by Cys as

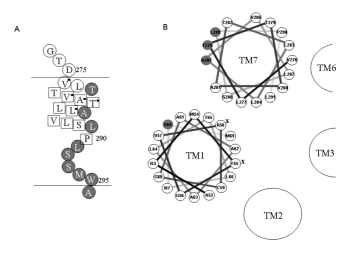


FIGURE 8: Representation of cross-linking results of TM7 and helical wheel presentation of TM1 and TM7 in Ste2p. (A) Summary of TM7 cysteine cross-linking results. The horizontal lines represent the boundary of the cell membrane. Residues indicated by a circle filled in gray when mutated to Cys are involved in dimer formation, whereas those residues shown in a box do not form disulfide bonds. Residues marked by an * compensate for the nonfunctional Y266C mutation. (B) Helical wheel presentation of TM1 and TM7. Residues indicated by a circle filled in gray are involved in dimer formation as found in this study. Residues marked by an X are important for ligand binding.

illustrated in the modified model (Figure 7B). Therefore, the dimerization mediated by T278, A285, and L289 is consistent with an α -helical periodicity in TM7 that continues up to Pro 290. In the absence of a crystal structure for Ste2p, the disulfide crosslinking results contribute to understanding structural features of the functional receptor such as interhelical interactions that may be involved in oligomerization.

In contrast to the specific pattern of residues which could participate in cross-linking in the portion of TM7 most adjacent to the extracellular surface, when the engineered cysteine residues were located at the intracellular end of the TM7, C-terminal to Pro290 which creates a kink in TM7, all residues were able to form disulfide bonds (Figure 8A) implying an absence of an ordered structure. We do not believe that the experimental protocol, specifically the use of the oxidizing agent, leads to a global denaturation of Ste2p because native cysteine residues in TM1 (C59) and TM5 (C252) did not form an inter-GPCR disulfide linkage under the same experimental conditions (Figure 2B). Additionally, only one of the TM1 cysteine mutants formed Ste2p dimers and only 3 residues out of 12 in the extracellular part of the TM7 showed increased dimerization. Rather, we propose that the L291–W295 region of Ste2p may be flexible enough to allow these residues to be exposed for disulfide bond formation with their counterpart in a second Ste2p monomer. Proline residues disrupt hydrogen bonding and have been shown to twist the standard structure of helices by introducing a kink between the segments contiguous with each other and to form molecular hinges (41, 63). The dynamic nature of the residues at the carboxyl side of Pro290 is consistent with the results of a high-resolution analysis of a 73-residue fragment of Ste2p in DPC micelles which indicated that the Pro290 resulted in a kinked helix in this membrane mimetic environment (64). An irregular helical structure in part of a TM has been documented in the crystal structures of other GPCRs. In rhodopsin, TM7 contains a helical segment near the extracellular face of the receptor, followed by a nonhelical segment in the vicinity of

P303 (36). The crystal structure of the beta-adrenergic receptor also indicates the presence of a short extended area in TM7 (65). In addition, it has been proposed, using computational simulations of the NPXXY motif, that this region was not an ideal alpha helical structure (66). Thus, there is precedent to propose that the part of TM7 in Ste2p that is near the cytoplasmic membrane interface is not helical in structure. Although there is a possibility that the residues C-terminal to Pro290 are not in the transmembrane domain, this is unlikely because a previous study (42) and our cysteine accessibility results (data not shown) for these residues, strongly suggested that W295 and A296 are the boundary between TM7 and the intracellular space. Obviously, non-hydrogen bonded elements of a peptide in a membrane represent a high energy state and it is possible that critical water molecules satisfy some of the required interactions for Ste2p. Water molecules that bridge various receptor regions have been implicated in the crystal structure of other GPCRs (36, 65). Their involvement in the structure of Ste2p must await a high resolution structure of this GPCR.

It has been observed that the homodimer interface of the dopamine D2 receptor can be altered upon ligand binding (50). To monitor possible ligand-induced conformational changes in Ste2p dimerization, membrane samples were incubated with either agonist or antagonist followed by Cu-P treatment. TM1 mutants showed similar degrees of dimer formation in the presence or absence of ligand, which is consistent with a recent study showing that dimer formation of V45C at TM1 was insensitive to α -factor treatment (21). In contrast, both agonist and antagonist treatment led to a decrease in the level of dimer formed by TM7-TM7 interaction indicating that ligand binding caused a conformational change in this region.

The effect of ligand binding on TM7-mediated dimer formation suggests that resting state dimers and activated state dimers involve different TM arrangements. Previously it was shown that the ratio of monomer to dimer did not change in the presence of ligand in Ste2p as tested by FRET (17). Thus, it is possible that the equilibrium distribution between monomer-dimer is maintained during receptor activation, even while the dimer interface may differ. In addition, a recent study of the serotonin 5HT2c receptor demonstrated that ligand binding induces a differential effect on the dimer interfaces. While the TM1/TM1 interface was insensitive to ligand treatment, the TM4/5 interface changed during receptor activation (67).

The similar effects of agonist and antagonist binding on disulfide formation involving residues in the TM7 dimer interface can be understood in terms of distinct binding and signaling domains for α -factor (68). Extensive structure—function analysis of α-factor and Ste2p has revealed that the C-terminus of α-factor is critical for receptor binding. Since agonist (WHWL-QLKPGQPNleY) and antagonist (WLQLKPGQPNleY) are identical except for the antagonist's N-terminal truncation, it is reasonable to suppose that the binding sites of both ligands in the receptor are highly overlapping especially at the C-terminal end of the ligands. Moreover, the carboxyl terminus has been shown to interact with residues in TM1 (23, 69). On the basis of this understanding, it is not unreasonable that both agonists and antagonists first bind to TM1 promoting similar changes in the conformation of both TM1 and of TM7.

Interestingly, a number mutations of residues at the extracellular regions of TM1 and TM7 were found to compensate for the activity of the signaling defective mutant receptor Y266C

(Figure 7A, TM7 residues labeled as *)(70) indicating these two domains are important for receptor activation. The decrease in dimer formation at TM7 after ligand binding that we showed for T278C and A285C, both located in the extracellular part of TM7, also suggest a conformational change in this portion of Ste2p. Therefore, the extracellular ends of TM1 and TM7 may play a key role in initiating receptor activation after ligand binding to TM1 and the subsequent conformational change of TM7 may facilitate the receptor to adopt a fully activated state. Many GPCRs assume different conformations in their active and inactive states, and it is widely accepted that GPCRs sample many states even when occupied by ligand [for review see ref 71]. Under our experimental conditions Cu-P mediated disulfide formation appears to distinguish these different conformations, possibly implicating a change of the dimer interface as part of the receptor activation mechanism. Nonetheless, we cannot rule out the possibility that the changes in the TM7 dimer interface upon ligand binding reflect structural changes elsewhere in Ste2p.

Much remains to be understood concerning the significance of GPCR dimerization in the signaling process. The residues in TM7 examined in this study, S²⁸⁸LPLSSMWA²⁹⁶, are highly conserved in Class D receptors and are considered to have a function similar to the NPXXY residues in Class A receptors (35). In Class A receptors, the NPXXY motif is regarded to be important for receptor activation (72-75). It is possible that TM7 plays an important role in both receptor activation and dimerization in Ste2p and in other GPCRs as well, and that dimerization is linked to signaling by these ubiquitous proteins. The results presented herein provide residue level information concerning the proximity of different sites on two Ste2p monomers in their native membrane environment. The physiological relevance of this information and of dimerization in GPCRs continues to be an intriguing area of GPCR biology.

ACKNOWLEDGMENT

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