Regulation of GPCR expression through an interaction with CCT7, a subunit of the CCT/TRiC complex

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ABSTRACT
Mechanisms that prevent aggregation and promote folding of nascent G protein-coupled receptors (GPCRs) remain poorly understood. We identified chaperonin containing TCP-1 subunit eta (CCT7) as an interacting partner of the β isoform of thromboxane A2 receptor (TPβ) by yeast two-hybrid screening. CCT7 coimmunoprecipitated with overexpressed TPβ and β2-adrenergic receptor (β2AR) in HEK 293 cells, but also with endogenous β2AR. CCT7 depletion by small interfering RNA reduced total and cell-surface expression of both receptors and caused redistribution of the receptors to juxtanuclear aggresomes, significantly more so for TPβ than β2AR. Interestingly, Hsp90 coimmunoprecipitated with β2AR but virtually not with TPβ, indicating that nascent GPCRs can adopt alternative folding pathways. In vitro pull-down assays showed that both receptors can interact directly with CCT7 through their third intracellular loops and C-termini. We demonstrate that Trp334 in the TPβ C-terminus is critical for the CCT7 interaction and plays an important role in TPβ maturation and cell-surface expression. Of note, introducing a tryptophan in the corresponding position of the TPα isoform confers the CCT7-binding and maturation properties of TPβ. We show that an interaction with a subunit of the CCT/TCP-1 ring complex (TRiC) chaperonin complex is involved in regulating aggregation of nascent GPCRs and in promoting their proper maturation and expression.

INTRODUCTION
G protein-coupled receptors (GPCRs) form the largest and one of the most-studied families of cell-surface proteins. They respond to a vast array of cellular mediators, including hormones, neurotransmitters, lipids, nucleotides, peptides, ions, and photons. GPCRs have one of the widest therapeutic ranges and were estimated to be the targets of more than 30% of all marketed drugs (Jacoby et al., 2006; Salon et al., 2011). To be functional, these receptors must be correctly folded and transported to the proper location, usually the plasma membrane, in order to be activated by their respective ligands. Their seven-transmembrane structure with an extracellular N-terminus, alternating intra- and extracellular loops, and an intracellular C-terminus renders folding of GPCRs a complex process (Tao and Conn, 2014). Failure to achieve correct folding results in their retrotranslocation, ubiquitination, and endoplasmic reticulum (ER)-associated degradation (Conn and Ulloa-Aguirre, 2011). Dysregulation of GPCR folding, trafficking, and signaling contributes to...
a number of pathophysiologic processes (Belmonte and Blaxall, 2011; Conn and Ulloa-Aguirre, 2011; Vassart and Costagliola, 2011; Lappano and Maggioni, 2012). Given the importance of these receptors, it is critical to understand the mechanisms that regulate their correct expression, folding, and export as nascent polypeptides, which, despite an increasing number of studies in this field of research, remain poorly characterized.

Secreted and membrane proteins possessing an N-terminal signal peptide are recognized by the signal recognition particle (SRP), leading to the cotranslational insertion of the polypeptides directly in the ER membrane through a translocon-dependent mechanism. Only 5−10% of known GPCRs contain a signal peptide that leads to their direct insertion into the ER membrane (Schülein et al., 2012). Subsequent folding, posttranslational modifications, and trafficking are controlled by ER-resident proteins and chaperones (Roux and Cottrell, 2014). However, little is known regarding what happens to the majority of GPCRs that do not contain signal sequences in their N-termini. Studies have shown that transmembrane segments of GPCRs can act as signal anchor (SA) sequences and be recognized by the SRP, but it remains unclear how and when such recognition occurs (Audigier et al., 1987; Schülein et al., 2012). Unlike the signal peptide, the SA is not cleaved after translocon-mediated insertion into the ER. Since translation of membrane proteins lacking a signal peptide begins in the cytosol, the SRP has a very short window of time to bind the translating ribosome and recognize the SA, because their interaction is inversely proportional to the polypeptide length (Berndt et al., 2009). If the SRP is unable to bind the SA, the synthesized protein is exposed to the cytosolic environment, which can result in aggregation and misfolding (White et al., 2010). To prevent this from happening, eukaryotic cells possess chaperone proteins that assist the folding process of nascent polypeptides, maintaining them in an intermediate state of folding competence for posttranslational translocation in subcellular compartments. Two complexes of chaperone proteins have been identified to interact posttranslationally with near nascent proteins and seem to affect their translocation into the ER. The first is the well-known 70-kDa heat shock protein (Hsp70) system, and the second is the tailless complex polypeptide 1 (TCP-1), a group II chaperonin, also known as the CCT/TRiC-1 ring complex (TRiC complex; Deshaies et al., 1988; Plath and Rapoport, 2000). The exact sequence of posttranslational events leading to ER insertion is not fully understood, but studies have proposed a three-step process. First, the nascent peptide emerging from ribosomes is able to interact with the nascent polypeptide-associated complex or the SRP, which both regulate translational flux (Kirstein-Miles et al., 2013). However, once translation is completed, these proteins are no longer able to bind the polypeptide. Second, Hsp70 and/or CCT/TRiC complexes bind polypeptides to maintain a translocable state by preventing premature folding, misfolding, and aggregation (Melville et al., 2003; Cuéllar et al., 2008). Third, ER-membrane insertion is mediated by the translocon, which strips away the cytosolic chaperones. This process is called the posttranslational translocation pathway (Ngosuwan et al., 2003).

CCT/TRiC is a large cytosolic chaperonin complex of ∼900 kDa composed of two hetero-oligomeric stacked rings able to interact with nascent polypeptides, which mediates protein folding in an ATP-dependent manner and prevents aggregation in eukaryotes (Knee et al., 2013). Each ring consists of eight different subunits (CCT1 to CCT8) that share ∼30% sequence homology, particularly in their equatorial domains, which mediate interactions between subunits (Valpuesta et al., 2002). CCT/TRiC was originally characterized for its role in the folding of α-actin (Llorca et al., 1999). In recent years, the list of identified substrates for this complex has grown in both number and variety. For example, it has been shown that CCT/TRiC can bind and/or help the folding of different WD-repeat proteins like the G-protein β subunits (Plimpton et al., 2015), STAT3 (Kasembi et al., 2014), the von Hippel-Lindau (VHL) protein (Melville et al., 2003), and even membrane proteins such as the peroxisome membrane protein Pmp22p (Pause et al., 1997) and the cell-surface receptor LOX-1 (Bakthavatsalam et al., 2014). The central cavity formed by the two stacked rings protects the newly synthesized polypeptide against aggregation in the crowded environment of the cytosol. This large cavity can accommodate a wide range of protein sizes from 20 to 200 kDa (Dekker et al., 2008). It was shown that CCT/TRiC can cotranslationally bind nascent polypeptides exposing at least 50 amino acids outside of the ribosome exit channel, but this number can vary between the different substrates (McCallum et al., 2000).

Here we report a direct and functional interaction between GPCRs and the chaperonin containing TCP-1 subunit eta (CCT7). We provide evidence that CCT7 is involved in preventing aggregation of GPCRs and in regulating their expression, maturation, and transport to the cell surface. To our knowledge, this is the first demonstration of a functional interaction between GPCRs and the CCT/TRiC complex.

RESULTS

CCT7 interacts with the β-isoinform of thromboxane A2 receptor (TPβ) and β2-adrenergic receptor (β2AR)

A yeast two-hybrid screen was performed using the C-terminus of TPβ as bait with a human HeLa cell MATCHMAKER cDNA library. Roughly 3 × 10⁵ clones were screened, resulting in more than 600 positives. Approximately 200 of these clones showed strong growth on selective yeast medium (Trp-, Leu-, His-, and Ade-) and were isolated, characterized by dideoxy sequencing, and then aligned using the NCBI BLAST alignment search tool. Five independent clones covering the full-length CCT7 coding sequence were identified in this screen. As shown in Figure 1A, only yeast transformed with the pAS2.1-TPβ C-terminus and pGAD-CCT7 showed strong growth on selective Trp-, Leu-, His-, and Ade- yeast medium, indicating that CCT7 interacts with the TPβ C-terminus.

Interestingly, we also identified CCT7 as a putative interactor of the β2AR using a gel-free proteomic approach, but the interaction and its functional consequences were not characterized (Roy et al., 2013). To assess the interaction between CCT7 and both GPCRs in a cellular context, we performed immunoprecipitation experiments in HEK 293 cells transfected with pcDNA3-FLAG-β2AR or pcDNA3-HA-TPβ and pcDNA3-CCT7-c-myc epitope (MYC). Cell lysates were incubated with FLAG-specific or hemagglutinin (HA)-specific monoclonal antibodies, and coimmunoprecipitation of CCT7 was detected by Western blot analysis using a MYC-specific antibody. CCT7 coimmunoprecipitated with both receptors (Figure 1, B and C), confirming that CCT7 forms a complex with either TPβ or β2AR in cells. Importantly, our data further show that the interaction can be detected endogenously. Indeed, HEK 293 cell lysates were incubated with CCT7-specific or nonspecific goat antibodies and coimmunoprecipitation of endogenous β2AR was detected by Western blot analysis using a β2AR-specific antibody. Endogenous β2AR was coimmunoprecipitated only when endogenous CCT7 was immunoprecipitated (Figure 1D).

Depletion of CCT7 impairs β2AR and TPβ total and cell-surface expression

To characterize the functional role of CCT7 interactions with β2AR and TPβ, we first investigated the effect of CCT7 depletion on total...
expression of the receptors by Western blot. HEK 293 cells stably expressing HA-β2AR or HA-TPβ were transfected with control or CCT7 dicer-substrate short interfering RNAs (DsiRNAs). Figure 2, A and B, shows that the partial depletion of CCT7 leads to decreased total protein expression of both receptors. Densitometry analyses of multiple independent experiments revealed that CCT7 depletion resulted in a loss of 42 and 37% in total receptor expression for TPβ and β2AR, respectively (Figure 2, C and D). We then assessed the importance of CCT7 expression on the cell-surface expression of β2AR, TPβ, and thromboxane A2 receptor α-isof orm (TPα), a shorter isoform of TP compared to TPβ generated by alternative splicing of the receptor C-terminus (Raychowdhury et al., 1994). HEK 293 cells expressing HA-tagged TPβ, TPα, or β2AR were transfected with control or CCT7 DsiRNAs and receptor cell-surface expression was measured by enzyme-linked immunosorbent assay (ELISA). CCT7 depletion significantly decreased TPβ cell-surface expression by 50% (Figure 2E), whereas cell-surface expression of TPα and β2AR was less affected, with reductions of 30 and 22%, respectively (Figure 2, F and G).

In mammals, nascent polypeptides emerging from ribosomes are bound by Hsc70, which operates in concert with the CCT/TRiC complex in the folding of polypeptides. Alternatively, polypeptides can be passed to Hsp90 (Young et al., 2004). In an attempt to address why CCT7 depletion affected the cell-surface expression of TPβ more significantly than that of β2AR, we compared the ability of both receptors to interact with Hsp90. Lysates of HEK 293 cells expressing FLAG-β2AR or FLAG-TPβ and HA-Hsp90 were immunoprecipitated with a Flag-specific antibody, and Hsp90 interaction was studied by Western blot analysis with an HA-specific monoclonal antibody (Figure 2H). Interestingly, significantly more Hsp90 protein coimmunoprecipitated with β2AR than with TPβ.

**CCT7 colocalizes with β2AR and TPβ and regulates their intracellular distribution**

Confocal microscopy was performed in HEK 293 cells stably expressing HA-β2AR or HA-TPβ that were transfected or not with control or CCT7 DsiRNAs and labeled with HA- and CCT7-specific antibodies. Figure 3, Ab and Bb, shows the levels of nonspecific secondary antibody–associated staining compared with the labeling obtained with the CCT7-specific antibody (Figure 3, Af and Bf). Endogenous CCT7 displayed cytosolic and nuclear localization in control DsiRNA-treated cells. HA-TPβ was distributed at the plasma membrane and in intracellular compartments in nontreated and control DsiRNA-transfected cells (Figure 3A, c and g), as we reported before (Thériault et al., 2004; Parent et al., 2009). Under the same conditions, HA-β2AR was primarily localized to the plasma membrane with some intracellular labeling (Figure 3B, c and g). Analysis of merged images revealed that CCT7 mainly colocalized with both receptors in the juxtanuclear region of the cell (Figure 3, Ah and Bh). Transfection of CCT7 DsiRNAs significantly diminished expression of endogenous CCT7 (Figure 3, Aj and Bj) and caused a marked redistribution of both receptors to an intracellular and juxtanuclear localization, which was more pronounced for HA-TPβ (Figure 3, Ak and Bk), in agreement with data obtained in receptor cell-surface expression experiments (Figure 2, E and G). Depletion of CCT7 also appeared to decrease receptor-associated fluorescence for both
receptors, also supporting our findings from Western blot analyses (Figure 2, A and B).

**CCT7 depletion induces accumulation of misfolded receptors in intracellular aggregates**

Because the distribution of the receptors was reminiscent of Golgi localization in cells transfected with CCT7 DsiRNAs, we performed colocalization studies between TPβ and GM130, a Golgi marker, in CCT7-depleted HEK 293 cells (Figure 4A). Partial colocalization was observed between the receptor and GM130 (Figure 4A). The colocalization of misfolded proteins to a juxtanuclear localization and a spatial overlap with the Golgi apparatus have been demonstrated to be associated with the formation of aggresomes (Johnston et al., 1998; García-Mata et al., 1999; Salemi et al., 2014). Aggresomes are made up of aggregated inclusion bodies and misfolded proteins (Watanabe et al., 2012). Given the function of CCT7 in protein folding, we reasoned that the receptors could be found in aggresomes in CCT7-depleted cells. Confocal microscopy was performed as above in HEK 293 cells stably expressing HA-TPβ or HA-β2AR transfected with control or CCT7 DsiRNAs (Figure 4, B and D). Cells were also stained with a probe, the PROTEOSTAT dye, designed to detect aggresomes by recognition of inclusion bodies and misfolded proteins. Low levels of colocalization were detected between the receptors and aggresomes under control conditions represented by low Manders’ colocalization coefficients of 0.03 and 0.01 for TPβ and β2AR, respectively (Figure 4, C and E). However, CCT7 depletion resulted in increased colocalization of both receptors with aggresomes in a juxtanuclear region (Figure 4, Bf and Df). This was more drastic for TPβ than for β2AR, as indicated by Manders’ colocalization coefficients in CCT7-depleted cells of 0.84 and 0.30 for TPβ and β2AR, respectively (Figure 4, C and E). These results indicate that CCT7 depletion induced an accumulation of misfolded TPβ and β2AR in intracellular aggregates, notably more pronounced for the former. It is also interesting to observe an overall augmentation of the aggresome staining across the cytosol of CCT7-depleted cells compared with the control (Figure 4, Be and De). This is probably caused by the detection, by the PROTEOSTAT dye, of other broadly distributed misfolded proteins.

**Determination of the CCT7-binding domain on β2AR and TPβ**

To establish whether or not the interaction of CCT7 with receptors could be direct, and if so to determine its binding domains on β2AR and TPβ, we performed in vitro binding assays with purified forms of recombinant intracellular loops (ICL) or C-termini (CT) of both receptors fused to glutathione S-transferase (GST) along with purified CCT7-MYC fused with a hexa-His tag (His6-CCT7-MYC). We also investigated whether CCT7 interacted with the C-terminus of TPα, a C-terminal spliced isoform if so to determine its binding domains on the receptor and GM130 (Figure 4Ad). The partial colocalization of both receptors with aggresomes under control conditions represented by low Mander’s colocalization coefficients of 0.03 and 0.01 for TPβ and β2AR, respectively (Figure 4, C and E). However, CCT7 depletion resulted in increased colocalization of both receptors with aggresomes in a juxtanuclear region (Figure 4, Bf and Df). This was more drastic for TPβ than for β2AR, as indicated by Manders’ colocalization coefficients in CCT7-depleted cells of 0.84 and 0.30 for TPβ and β2AR, respectively (Figure 4, C and E). These results indicate that CCT7 depletion induced an accumulation of misfolded TPβ and β2AR in intracellular aggregates, notably more pronounced for the former. It is also interesting to observe an overall augmentation of the aggresome staining across the cytosol of CCT7-depleted cells compared with the control (Figure 4, Be and De). This is probably caused by the detection, by the PROTEOSTAT dye, of other broadly distributed misfolded proteins.

**FIGURE 2: CCT7 depletion impairs TPβ and β2AR total and cell-surface expression.** HEK 293 cells stably expressing HA-TPβ (A) or HA-β2AR (B) were transfected with control DsiRNA (DsiCtrl) or CCT7 DsiRNA (DsiCCT7), and lysates were immunoblotted with HA-specific HRP-conjugated antibodies. Densitometry was performed on the Western blots to quantify relative expression of HA-TPβ (C) and HA-β2AR (D) in cells treated with CCT7 DsiRNA compared with control DsiRNA-transfected cells (100%) and normalized to GAPDH expression. Densitometry was performed using ImageJ software, and the results are presented as mean ± SD of at least four independent experiments. Cell-surface receptor expression was measured in HEK 293 cells expressing HA-TPβ (E), HA-TPα (F), or HA-β2AR (G) transfected with control or CCT7 DsiRNAs by ELISA using a monoclonal HA-specific antibody as described in Materials and Methods. Results are shown as a percentage of receptor expression measured in HEK 293 cells stably expressing HA-TPβ or HA-Hsp90 alone or together were immunoprecipitated with FLAG-specific polyclonal and GAPDH-specific antibodies. Densitometry was performed on the Western blots to quantify relative expression of HA-TPβ and HA-TPα (H) DsiRNAs (Figure 4, B and D). Cells were also stained with a probe, the PROTEOSTAT dye, designed to detect aggresomes by recognition of inclusion bodies and misfolded proteins. Low levels of colocalization were detected between the receptors and aggresomes under control conditions represented by low Manders’ colocalization coefficients of 0.03 and 0.01 for TPβ and β2AR, respectively (Figure 4, C and E). However, CCT7 depletion resulted in increased colocalization of both receptors with aggresomes in a juxtanuclear region (Figure 4, Bf and Df). This was more drastic for TPβ than for β2AR, as indicated by Manders’ colocalization coefficients in CCT7-depleted cells of 0.84 and 0.30 for TPβ and β2AR, respectively (Figure 4, C and E). These results indicate that CCT7 depletion induced an accumulation of misfolded TPβ and β2AR in intracellular aggregates, notably more pronounced for the former. It is also interesting to observe an overall augmentation of the aggresome staining across the cytosol of CCT7-depleted cells compared with the control (Figure 4, Be and De). This is probably caused by the detection, by the PROTEOSTAT dye, of other broadly distributed misfolded proteins.
role of Trp334 in TPβ maturation

Closer observation by Western blot analysis of the migration patterns of the TPβ W334Q and TPα Q333W mutants from cell lysates revealed that they were different from their respective wild-type counterparts. Figure 7A shows that wild-type HA-TPβ was detected as a major form at ~40 kDa and a minor form at ~70 kDa. In contrast, HA-TPβ W334Q displayed the opposite pattern, with minor and major forms at ~40 and ~70 kDa, respectively. The latter was reminiscent of the expression pattern shown by wild-type HA-TPα (Figure 7B), which migrated as a prominent form in the ~50–55 kDa range and as a minor form at ~35 kDa. On the other hand, the TPα Q333W mutation induced a redistribution of the higher molecular form of the receptor toward the lower ~35 kDa form, analogous to the distribution of the wild-type HA-TPβ.

We previously reported that the TPβ ~70 kDa and TPα ~50–55 kDa forms corresponded to glycosylated dimers of the receptors measured using treatment with peptide N-glycosidase F, which removes all types of N-linked oligosaccharides from glycoproteins (Laroche et al., 2005). Here we further analyzed the glycosylation patterns of TP isoforms with endoglycosidase H (Endo Hf), an N-glycosidase that selectively removes unprocessed high mannosetype N-linked oligosaccharides present on ER-resident glycoproteins. Glycosylated TP receptor proteins that have undergone trimming in the Golgi will be resistant to Endo Hf treatment. Lysates of HEK 293 cells expressing HA-TPβ or HA-TPα were treated with Endo Hf and then analyzed by Western blot. As shown in Figure 7C, the larger HA-TPβ ~70 kDa and TPα ~50–55 kDa forms were predominantly unaltered, whereas the lower forms of the receptors were reduced in size upon Endo Hf treatment. Altogether our present data, along with our earlier results (Laroche et al., 2005), indicate that the lower molecular weight bands of TPβ and TPα are immature monomeric forms of the receptors present in the ER. On the other hand, the higher molecular weight Endo Hf-resistant forms represent dimeric TP receptors that have undergone complex glycosylation in the Golgi. Consistent with this, our results suggest that the HA-TPβ W334Q mutation promoted receptor maturation through the Golgi toward a glycosylated receptor dimer (Figure 7A). In contrast, the
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Using ELISAs in HEK 293 cells, we then evaluated whether the effects of the TPβ W334Q and TPβ Q333W substitutions on receptor maturation were reflected in receptor cell-surface expression. Compared with wild-type HA-TPβ, cell-surface expression of HA-TPβ W334Q was increased by more than 200% to reach similar levels as HA-TPα (Figure 7D). Conversely, HA-TPα Q333W detection at the cell membrane was reduced by 50% compared with wild-type HA-TPα (115 vs. 239%), akin to wild-type TPβ expression levels. The effect of the TPβ W334Q mutation on cell-surface expression of the receptor was also studied by confocal microscopy. HEK 293 cells transiently expressing HA-TPβ or HA-TPβ W334Q were labeled with an HA-specific antibody to visualize the receptors and with a CCT7-specific antibody to detect endogenous CCT7 (Figure 8A). As we reported before, wild-type TPβ exhibited plasma membrane staining accompanied by strong intracellular localization (Figure 8Ac). On the other hand, the TPβ W334Q mutant displayed robust membrane localization (Figure 7Ag). Quantification of receptor immunofluorescence was carried out on 100 cells for each receptor construct. Figure 8B shows that 25% of wild-type TPβ immunofluorescence was found at the plasma membrane, compared with 55% for the TPβ W334Q mutant, a roughly twofold difference, confirming our cell-surface expression data obtained by ELISA (Figure 7D). We also observed that TPβ colocalized more significantly with CCT7 than did the TPβ W334Q mutant (Figure 8A, d and h). Quantification of CCT7 colocalization with the two receptor constructs revealed Mander’s colocalization coefficients of 0.43 for TPβ and 0.12 for TPβ W334Q (Figure 8C). This marked decrease in CCT7 colocalization with the TPβ W334Q mutant is in line with the virtual lack of detectable coimmunoprecipitation between the two proteins (Figure 6C).

Next we assessed the effect of CCT7 depletion on the colocalization of the TPβ W334Q mutant with the aggresome. Confocal microscopy experiments showed that the receptor mutant, in the presence of CCT7 DsiRNAs, was readily detected at the cell surface (Figure 9Ab) but also redistributed to the aggresome (Figure 9Af). Quantification of the colocalization between the TPβ W334Q mutant and the aggresome yielded a Mander’s coefficient of colocalization of 0.51 when CCT7 was depleted compared with 0.10 when the cells were transfected with control DsiRNAs (Figure 9B). These results indicate that the receptor mutant is still affected by CCT7 depletion in terms of redistribution to the aggresome but in a less drastic way than was wild-type TPβ (Figure 4, B and C, Mander’s coefficient of 0.84). Taken together, our results indicate that Trp334 of TPβ interacts with CCT7 and plays an important role in the maturation and cell-surface expression of the receptor.

CCT7 interacts with other GPCRs

Finally, knowing that CCT7 can interact with the β2AR and TPβ but not with TPα, we wanted to extend our coimmunoprecipitation studies to a few other GPCRs. Lysates of HEK 293 cells transiently expressing HA-tagged rat μ-opioid receptor (MOR; Figure 10A), FLAG-tagged rat δ-opioid receptor (DOR; Figure 10B), or FLAG-tagged prostaglandin D2 receptor (DP; Figure 10C) with or without CCT7-MYC were incubated with HA- or FLAG-specific antibodies. Coimmunoprecipitation of CCT7 was detected by Western blot analysis using a Myc-specific antibody. Coimmunoprecipitation of CCT7 was observed with each of the three receptors. Our data suggest that CCT7 can interact with various GPCRs, but not all, and that...
analysis of the CCT/TRiC substrates noted that the CCT/TRiC interactome is enriched in large, hydrophobic proteins with intricate topologies predicted to be slow folding and aggregation prone and belonging to oligomeric protein complexes. On the other hand, CCT/TRiC could fold monomeric subunits and maintain them in an assembly-competent state until association with oligomeric partners (Yam et al., 2008). Here we provide evidence that the CCT/TRiC chaperonins prevent aggregation and participate in the folding and maturation of the complex seven-transmembrane GPCR proteins known to form oligomeric structures between receptor units and with signaling partners during their biogenesis (Dupré and Hébert, 2006; Dupré et al., 2006).

We first identified the interaction between CCT7 and TPβ by yeast two-hybrid screening with the C-terminus of the receptor. We also detected all of the eight CCT subunits as potential β2AR-binding partners in liquid chromatography–tandem mass spectrometry proteomics studies (Roy et al., 2013). Our data showed that TPβ and β2AR can interact directly with CCT7 through their third intracellular loops and their C-termini. The interaction was confirmed by coimmunoprecipitation experiments in a cellular context in overexpression settings for both receptors but notably also between endogenous CCT7 and β2AR proteins. Further analysis also showed that CCT7 coimmunoprecipitated in HEK 293 cells with other GPCRs, such as the prostaglandin D2 DP1, and the μ- and δ-opioid receptors, suggesting that CCT7 may be involved in a general mechanism of GPCR biogenesis.

TPβ displays prominent intracellular localization in our system, which is consistent with the fact that it is prone to ER retention and ER-associated degradation as shown previously by us and others (Parent et al., 2008; Ball et al., 2010). Depletion of CCT7 resulted in lower total and cell-surface expression for both TPβ and β2AR. It also caused redistribution of the receptors to aggresomes, drastically in the case of TPβ. This indicates that CCT7 is involved in preventing aggregation and degradation of the receptors. Of note, it was shown that mutation of CCT7 in yeast caused the VHL protein to accumulate in juxtanuclear aggregates (Amit et al., 2010), similar to what we observed for the receptors in our study. It is also interesting that the misfolded GPCR rhodopsin P23H mutant, associated with autosomal-dominant retinitis pigmentosa, also accumulates in aggresomes and is ubiquitinylated and degraded by the proteasome (Saliba et al., 2002), further supporting the notion that inappropriately folded GPCRs are targeted to aggresomes. The receptors in CCT7-depleted cells colocalized with a subfraction of the misfolded proteins stained by the PROTEOSTAT aggresome dye, with a juxtanuclear localization that overlapped partially with the Golgi. Aggregated inclusion bodies and misfolded proteins accumulate in aggresomes (Watanabe et al., 2012). This aggregation process is more likely to occur cotranslationally while hundreds of nascent polypeptide chains emerge from the polysomes at the same time in a single localization leading to the improper folding (Garcia-Mata et al., 2002). The aggresomal particles are then transported toward the aggresome localization via dynein to be sequestered in a large single structure called the aggresome (Garcia-Mata et al., 1999). This juxtanuclear sequestration is a nontoxic cellular response preventing aggregation and degradation of the receptors. Of note, it was shown that mutation of CCT7 in yeast caused the VHL protein to accumulate in juxtanuclear aggregates (Amit et al., 2010), similar to what we observed for the receptors in our study. It is also interesting that the misfolded GPCR rhodopsin P23H mutant, associated with autosomal-dominant retinitis pigmentosa, also accumulates in aggresomes and is ubiquitinylated and degraded by the proteasome (Saliba et al., 2002), further supporting the notion that inappropriately folded GPCRs are targeted to aggresomes. The receptors in CCT7-depleted cells colocalized with a subfraction of the misfolded proteins stained by the PROTEOSTAT aggresome dye, with a juxtanuclear localization that overlapped partially with the Golgi. Aggregated inclusion bodies and misfolded proteins accumulate in aggresomes (Watanabe et al., 2012). This aggregation process is more likely to occur cotranslationally while hundreds of nascent polypeptide chains emerge from the polysomes at the same time in a single localization leading to the improper folding (Garcia-Mata et al., 2002).
in the folding and trafficking of other GPCRs, such as the melanocortin-4 receptor (Meimaridou et al., 2011), the prostaglandin D_2 receptor (Binda et al., 2014), the adenosine A_2A receptor (Bergmayr et al., 2013), and the α_2-adrenergic receptor (Filipeanu et al., 2011).

Our results demonstrated that Trp^{334} is crucial for TPβ to bind to CCT7 and that introduction of a tryptophan residue in the TPα C-terminus promoted its interaction with CCT7. Strikingly, the TPβ W334Q and TPα Q333W substitutions conferred properties that corresponded to the other wild-type TP isof orm. It has been shown that hydrophobic interactions are involved in the binding of CCT subunits to actin, VHL, and Gβ proteins (Kabir et al., 2011). In particular, replacement of Trp^{117} substantially reduced the binding of the VHL protein to CCT (Feldman et al., 2003), similar to what we observed for TPβ. It may be that CCT-complex proteins serve to facilitate early interactions between receptors and Gβ proteins during their biosynthesis and favor their proper assembly (Dupré et al., 2006). However, the fact that we identified a single residue, Trp^{334}, that dictates the interaction between CCT7 and TPα or TPβ and their maturation and trafficking properties strongly supports that CCT7 acts, at least in part, directly on the receptors.

It might be counterintuitive that TPβ Trp^{334} is a major determinant of the TPβ-CCT7 interaction but its mutation does not reduce total TPβ protein expression and promotes maturation and cell-surface expression of the receptor. It has been shown that depletion of a single CCT subunit leads to the codepletion of the other CCT/TRiC complex subunits (Freund et al., 2014). Thus expression knockdown of CCT7 could destabilize or deplete the CCT/TRiC complex, leading to receptor aggregation and degradation. Furthermore, it is common for substrates using the chaperonin complex to interact with multiple subunits through distinct determinants (Zhuravleva and Radford, 2014). It is reasonable to speculate that GPCRs using the CCT/TRiC complex will interact with multiple domains of many CCT subunits. This could explain, in part, why TPβ W334Q displays decreased CCT7 binding but is still able to mature and fold correctly, perhaps in the folding and trafficking of other GPCRs, such as the melanocortin-4 receptor (Meimaridou et al., 2011), the prostaglandin D_2 receptor (Binda et al., 2014), the adenosine A_2A receptor (Bergmayr et al., 2013), and the α_2-adrenergic receptor (Filipeanu et al., 2011).

The β_2AR was less affected than TPβ by CCT7 expression knockdown. This suggested that the β_2AR might take advantage of complementary folding pathways. Indeed, as nascent polypeptide chains emerge from ribosomes they are met by Hsc70. The latter may act together with CCT/TRiC in the folding of different polypeptides. Other newly synthesized polypeptides can fold spontaneously or be assisted by Hsc70. Alternatively, they can be passed endowed with Hsc70 to Hsp90 for subsequent folding (Young et al., 2004). Our data showed that Hsp90 coimmunoprecipitated with the β_2AR, whereas its interaction with TPβ was negligible. This indicates that nascent GPCR polypeptides can be subjected to distinct folding pathways depending on the receptor and could explain, at least in part, why the β_2AR cell-surface expression was less sensitive to CCT7 depletion than TPβ. Involvement of Hsp90 has been reported...
physiological concentrations. High- and low-affinity CCT subunits were shown to be spatially segregated within two contiguous hemispheres in the CCT/TRiC ring, generating an asymmetric power stroke that is thought to drive the folding cycle. This uncommon mode of ATP usage is thought to serve in coordinating a directional mechanism underlying the unique ability of CCT/TRiC to fold complex proteins (Reissmann et al., 2012). Further experiments will be needed to fully understand the specificity and the role of the various CCT subunits and their binding determinants involved in GPCR folding.

In summary, we have characterized a direct and functional interaction between a subunit of the CCT/TRiC chaperonin complex and GPCRs. We have provided evidence that distinct nascent GPCRs can undergo alternative folding pathways and that CCT/TRiC is critical in preventing aggregation of some GPCRs and in promoting their proper maturation and expression.

MATERIALS AND METHODS

Reagents

Monoclonal anti-HA (3F10) antibody was from Roche Applied Science (Pleasanton, CA). Anti-GAPDH polyclonal antibody, anti–TCP-1η (CCT7) (E-20) goat polyclonal antibody, anti–β2AR (H-73) polyclonal antibody, and protein G–agarose beads were purchased from Santa Cruz Biotechnology (Santa Cruz, CA). Texas Red anti-goat immunoglobulin G (IgG) was from Jackson ImmunoResearch (West Grove, PA). Alexa Fluor 488, Alexa Fluor 633 anti-rabbit IgG, and ProLong Gold antifade reagent were purchased from Invitrogen (Carlsbad, CA). Anti–c-Myc (9E10) monoclonal antibody was from Covance (Montreal, Canada). Anti–Myc-horseradish (HRP) polyclonal antibody was from Abcam (Toronto, Canada). Anti–rabbit-HRP IgG and anti–mouse-HRP IgG were purchased from GE Healthcare (Baie d’Urfé, Canada). Anti–goat-HRP IgG was from R&D Systems (Minneapolis, MN). Polyclonal anti–FLAG, the alkaline phosphatase–conjugated anti-mouse antibodies, and the alkaline phosphatase substrate kit were purchased from Sigma-Aldrich (Oakville, Canada). PROTEOSTAT Aggresome Detection Kit was purchased from BioVision (Milpitas, CA).
from Enzo Life Sciences (Farmingdale, NY). Goat anti–GST-HRP antibody was purchased from Bethyl Laboratories (Montgomery, TX). Mouse anti-HA IgG, and stained with PROTEOSTAT aggresome dye. Alexa Fluor 633–conjugated anti-mouse IgG was used as the secondary antibody. The third images represent a merged image (c and f) of the green and red signals where the areas with high degree of colocalization between the green signal of the receptors (a and d) and red signal of the aggresome (b and e) appear yellow. Scale bars: 10 μm. Images shown are single confocal slices representative of at least four independent experiments and more than 250 observed cells. Mander’s colocalization coefficients represent the ratio of the green signal of the receptor overlapping the red signal of aggresomes and were calculated from at least 100 cells per condition. Results are presented as mean ± SEM.

FIGURE 9: Targeting of the TPβ Trp334 mutant to the aggresome is diminished compared with wild-type TPβ in CCT7-depleted cells. (A) HEK 293 cells transiently expressing HA-TPβ W334Q were treated with control or CCT7 DsiRNAs. The cells were fixed, permeabilized, labeled with mouse anti-HA IgG, and stained with PROTEOSTAT aggresome dye. Alexa Fluor 633–conjugated anti-mouse IgG was used as the secondary antibody. The third images represent a merged image (c and f) of the green and red signals where the areas with high degree of colocalization between the green signal of the receptors (a and d) and red signal of the aggresome (b and e) appear yellow. Scale bars: 10 μm. Images shown are single confocal slices representative of at least four independent experiments and more than 250 observed cells. (B) Mander’s colocalization coefficients represent the ratio of the green signal of the receptor overlapping the red signal of aggresomes and were calculated from at least 100 cells per condition. Results are presented as mean ± SEM.

FIGURE 10: CCT7 coimmunoprecipitates with other GPCRs. (A) Lysates of HEK 293 cells transiently expressing HA-MOR (HA-tagged rat μ-opioid receptor) alone or with CCT7-MYC were immunoprecipitated with an HA-specific monoclonal antibody and analyzed by immunoblotting with MYC- and HA-specific HRP-conjugated antibodies. Lysates of HEK 293 cells transiently expressing FLAG-DOR (FLAG-tagged rat δ-opioid receptor; B) or FLAG-DP (FLAG-tagged prostaglandin D2 receptor; C) alone or with CCT7-MYC were immunoprecipitated with a FLAG-specific monoclonal antibody and analyzed by immunoblotting with FLAG-specific polyclonal and HA-specific HRP-conjugated antibodies. The blots shown are representative of three separate experiments. IB, immunoblotting; IP, immunoprecipitation.

Yeast two-hybrid screen
A yeast two-hybrid screen was performed following the two-hybrid system standard protocol (Gietz and Woods, 2002). Briefly, a plasmid containing the C-terminus of TPβ (pAS2-1-TPβCT) was transformed into the yeast strain pJ69-4αx using the lithium yeast transformation protocol (Gietz and Woods, 2002). This stably transformed clone was then transformed with a human HeLa cell MATCHMAKER cDNA Library or with the empty pGAD-424 plasmid (Clontech, Mountain View, CA). Positive clones were initially selected for growth in the absence of histidine, and interactions were confirmed by growth on quadruple-selective medium (Trp-, Leu-, His-, and Ade-) and transformed into the DH10B bacterial strain. Plasmids were extracted from DH10B cells and transformed once more into yeast with either the bait (pAS2-1-TPβCT) or the negative control (pAS2-1) and plated on quadruple-selective medium (Trp-, Leu-, His-, and Ade-) to confirm the interaction. The selected plasmids were then sequenced by dideoxy DNA sequencing, and the identities of the clones were determined by using the NCBI BLAST alignment tool.

Cell culture and transfection
Human embryonic kidney 293 (HEK 293) cells were maintained in DMEM (Invitrogen) supplemented with 10% fetal bovine serum at 37°C in a humidified atmosphere containing 5% CO2. Transient transfection of HEK 293 cells grown to 50–70% confluence was performed using the TransIT-LT1 Reagent (Mirus, Madison, WI) according to the manufacturer’s instructions. Empty pcDNA3 vector was added to keep the total DNA amount constant per plate. Stably TPβ- and β2AR-expressing HEK 293 cells were generated as previously described (Azzi et al., 2003; Parent et al., 2008) and cultured the same way as transiently transfected cells except for the addition of 200 μg/ml of G418.

The synthetic duplex oligonucleotide named HSC.RNAI. N006429.12.4 targeting the human CCT7 gene and the negative control DsiRNA (DS NC1, catalogue number- 51-01-14-03) were
purchased from Integrated DNA Technologies (Coralville, IA). HEK 293 cells were transfected with 50 nM oligonucleotides using the Lipofectamine 2000 transfection reagent (Invitrogen) according to the manufacturer’s recommendations, except for the following modifica- tions: Cells were seeded directly into the transfection mix at twice the cell density indicated in the basic protocol. Reverse tran- scriptase-PCRs were carried out to confirm that the CCT7 DsiRNAs did not reduce the mRNA levels of the receptors.

**Immunoprecipitations**

HEK 293 cells were transiently transfected with the indicated con- structs and maintained as described above for 48 h. Cells were then washed with ice-cold phosphate-buffered saline (PBS) and har- vested in 300 μl of lysis buffer (150 mM NaCl, 50 mM Tris, pH 8.0, 0.5% deoxycholate, 0.1% SDS, 10 mM Na3P2O7, 1% IGEPAL, and 5 mM ethylenediaminetetraacetic acid) supplemented with prote- ase inhibitors (10 μM pepstatin, 10 μM antipain, 10 μM leupeptin, and 10 μM chymostatin [Sigma-Aldrich]). After 60 min of incubation in lysis buffer at 4°C with rotation, the lysates were then centrifuged for 20 min at 14,000 × g at 4°C. One microgram of specific antibod- ies was added to the supernatant. After 3 h of incubation at 4°C with rotation, 40 μl of 50% protein G–agarose beads was added, fol- lowed by overnight incubation at 4°C. Samples were then centri- fuged for 1 min in a microcentrifuge and washed four times with 1 ml of lysis buffer. Immunoprecipitated proteins were eluted by addition of 35 μl of SDS sample buffer, followed by a 60 min incuba- tion at room temperature. Initial lysates and immunoprecipitated proteins were analyzed by SDS–PAGE and immunoblotting with specific antibodies. Endogenous immunoprecipitations were per- formed in native HEK 293 cells. Cells were harvested and processed as described above, except proteins were immunoprecipitated overnight using 2 μg TCP-1p (CCT7)-specific or appropriate control antibodies and 40 μl of 50% protein G–agarose beads.

**Immunofluorescence staining and confocal microscopy**

For colocalization experiments, HEK 293 cells stably expressing HA-β2AR or HA-TPβ were plated in six-well plates at a density of 5 × 10⁴ cells/well directly onto coverslips coated with 0.1 mg/ml poly-l-lysine (Sigma-Aldrich) and transfected with control or CCT7-specific DsiRNAs. The cells were fixed after a 72 h incubation with 2% (vol/vol) formaldehyde/PBS at 30 min at 4°C. Subsequently cells were washed twice with PBS and permeabilized for 10 min with 0.1% Triton X-100 in PBS and blocked for 30 min with 0.1% Triton X-100 in PBS containing 0.5% (wt/vol) bovine serum albumin (BSA) at room temperature. After two washing steps with 0.1% Triton X-100 in PBS, cells were incubated 2 h with HA-specific and CCT7- specific (not for IgG Ctrl conditions) antibodies diluted in blocking buffer at room temperature. The cells were washed twice with per- meabilization buffer, blocked again for 10 min, and incubated with appropriate secondary antibodies for 60 min at room temperature or with the Proteostat aggresome dye according to the manufac- turer’s recommendations. Cells were then washed three times with PBS, and coverslips were mounted using ProLong Gold antifade reagent. Confocal microscopy was performed using a scanning confocal microscope (FV1000; Olympus, Richmond Hill, Canada) coupled to an inverted microscope with a 60x oil-immersion objective (Olympus), and all laser parameters were conserved between all im- age acquisitions for the same figure. Images were processed using Fluoviewer 2.0 software (Olympus), and Mander’s colocalization co- efficients (Dunn et al., 2011) were calculated using the same thresh- old for fluorescent background elimination across all images, since they were acquired with the same parameters.

**Measurement of cell-surface receptor expression by ELISA**

Quantification of cell-surface receptor expression was carried out as described before (Binda et al., 2014). Briefly, 5 × 10⁴ HEK 293 cells stably expressing HA-β2AR or HA-TPβ were plated in 24-well plates precoated with 0.1 mg/ml poly-l-lysine (Sigma- Aldrich). Cells were transfected with the indicated DsiRNAs and then maintained for an additional 72 h. The cells were fixed in 3.7% (vol/vol) formaldehyde/Tris-buffered saline (TBS; 20 mM Tris-HCl, pH 7.5, and 150 mM NaCl) for 5 min at room temperature. Cells were then washed twice with TBS, and nonspecific binding was blocked by incubation with TBS containing 1% BSA for 30 min. A monoclonal HA-specific antibody was then added at a dilution of 1:2000 in TBS–BSA (1%) for 60 min. Following incubation with the primary antibody, cells were washed twice and blocked again with TBS–BSA (1%) for 10 min. Cells were then incubated with an alka- line phosphatase–conjugated goat anti-mouse antibody at 1:10,000 dilution in TBS–BSA (1%) for 60 min. Cells were washed twice with TBS, and 250 μl of a colorimetric alkaline phosphatase substrate was added as per the manufacturer’s instructions. The plates were then incubated at 37°C until a yellow color appeared. The reaction was stopped by the addition of 250 μl of NaOH (0.4 M). A 200 μl aliquot of the colorimetric reaction was taken, and the absorbance was measured at 405 nm using a TiterTek Multiskan MCC/340 spectrophotometer. All conditions were done in triplicate for each experiment.

**Recombinant protein production and histidine pull-down analysis**

For production of His-tagged proteins, a PCR fragment corre- sponding to the cDNA coding for full-length CCT7 was inserted into the pGEX-4T1 expression vector (Invitrogen) as described above. This construct was used to produce the fusion protein in OverExpressTM C41 (DE3) Escherichia coli strain (Avidis, Roubais, France) by following the manufacturer’s instructions. The recombinant proteins were purified using nickel–nitrilotriacetic acid–aga- rose resin (Qiagen, Toronto, Canada) as indicated by the manufac- turer. The cDNA fragments coding for the C-terminus and intracellular loops of β2AR or TPβ introduced in the pGEXT-4T1 vector (Amersham Biosciences, Baie d’Urfé, Canada) were used to produce GST fusion proteins in the OverExpressTM C41 (DE3) E. coli strain, which were purified using glutathione–Sepharose 4B beads (Amersham Biosciences) and eluted according to the manufac- turer’s indications. Purified recombinant proteins were analyzed by SDS–PAGE followed by Coomassie brilliant blue R-250 staining. Five micrograms of each GST-tagged fusion protein was incubated with 5 μg of the purified nickel–nitrilotriacetic acid–agarose His-tagged CCT7 in wash buffer (300 mM NaCl, 50 mM NaH2PO4, 20 mM imidazole, 0.5% Triton X-100, and 2 mM dithiothreitol) supplemented with protease inhibitors (9 nM pepstatin, 9 nM antipain, 10 nM leupeptin, and 10 nM chymostatin). Binding reac- tions were then washed three times with wash buffer. SDS sample buffer was added to the binding reactions, and the tubes were boiled for 5 min. The pull-down reactions were analyzed by West- ern blotting with the indicated specific antibodies.

**Deglycosylation assays**

Receptors were transiently expressed in HEK 293 cells in 60 mm plates, and cell lysates were prepared as described above. Protein concentration of samples was determined with the DC (deter- gent-compatible) Bio-Rad (St-Laurent, Canada) protein assay. For Endo H experiments, volumes of lysates corresponding to 40 μg protein were denatured in glycoprotein denaturing buffer 1×

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(New England Biolabs) for 30 min at 37°C followed by the addition of glycerol buffer 1× (New England Biolabs) and 10,000 U/ml of Endo Hf (New England Biolabs). Samples were incubated with the enzyme for 5 h at 37°C and analyzed by Western blot.

Statistical analysis
Statistical analysis was performed using Prism version 5.0 (GraphPad Software) using a two-tailed Student’s t test or one-way analysis of variance with multiple comparisons. Data were considered significant when p values were < 0.05 (*), < 0.01(**), < 0.001 (***) or < 0.0001 (****). Nonsignificant values are abbreviated as “ns.”

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