Identification and Characterization of Distinct C-Terminal Domains of the Human Hydroxycarboxylic Acid Receptor-2 That Are Essential for Receptor Export, Constitutive Activity, Desensitization, and Internalization

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ABSTRACT

The human hydroxycarboxylic acid receptor 2 (HCA2), also known as GPR109A and HM74a, was first identified as a niacin receptor and has recently received significant attention because of its potential to clinically modify plasma lipids in a favorable manner. Our recent studies have demonstrated that the niacin-induced internalization of HCA2 receptors is regulated by G protein-coupled receptor kinase (GRK) 2 and arrestin3 and that internalized receptors rapidly recycle back to the cell surface. The investigation presented here used a combination of amino acid deletion and site-directed mutagenesis to identify structural and functional domains within the HCA2 C terminus and explore their potential roles in receptor phosphorylation, desensitization, and internalization. We first constructed four mutants with deletions of 10 to 15 amino acids each that were distinct from truncated mutants. We successfully identified different domains responsible for receptor export, constitutive activity, desensitization, phosphorylation, and internalization. We also generated a comprehensive series of alanine substitution mutants, replacing conserved serine and threonine residues in the C terminus with alanine residues to pinpoint the key residues that are essential for GRK2-mediated phosphorylation and arrestin3 association. Moreover, we found that a sequence from residues 329 to 343 in the C-terminal tail of HCA2 plays a crucial role in keeping HCA2 in an inactive conformation. These data demonstrate the importance of distinct domains within the C terminus of HCA2 for receptor cell surface expression, desensitization, and internalization and phosphorylation and stabilization of an inactive receptor conformation.

Introduction

The high-affinity human G protein-coupled receptor for niacin (nicotinic acid), hydroxycarboxylic acid receptor 2 (HCA2), also known as GPR109A or HM74a, has drawn significant attention (Soga et al., 2003; Tunaru et al., 2003; Wise et al., 2003). HCA2 receptor activation not only lowers total plasma cholesterol, free fatty acid, and triglyceride levels but also significantly increases high-density lipoprotein cholesterol levels and thus offers a clinical treatment for dyslipidemia and atherosclerosis (Mack et al., 1993; Taylor et al., 1999). This work was supported by the Ministry of Science and Technology [Grants 2012CB910402, 2012AA020303-05]; National Natural Science Foundation of China [Grant 81173108]; Zhejiang Natural Science Foundation [Grant Z2080207]; and National Institutes of Health National Institute of General Medical Sciences [Grant GM44944].

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ABBREVIATIONS: GRK, G protein-coupled receptor kinase; ERK1/2, extracellular signal-regulated kinase1/2; siRNA, small interfering RNA; GPCR, G protein-coupled receptor; DiI, 1,1'-dioctadecyl-3,3,3',3'-tetramethylindocarbocyanine perchlorate; FITC, fluorescein isothiocyanate; HRP, horseradish peroxidase; HEK, human embryonic kidney; EGFP, enhanced green fluorescent protein; ELISA, enzyme-linked immunosorbent assay; WT, wild-type; TBS, Tris-buffered saline; BSA, bovine serum albumin; DMEM, Dulbecco’s modified Eagle’s medium; PBS, phosphate-buffered saline; CRE, cAMP response element; ER, endoplasmic reticulum; MjCD, methyl-β-cyclodextrin; PKA, protein kinase A; PKC, protein kinase C; 5-HT, 5-hydroxytryptamine.
2004). In 2003, the human orphan G protein-coupled receptor, HM74a, was identified as a niacin receptor (Soga et al., 2003; Tunaru et al., 2003; Wise et al., 2003). Two years later, the ketone body, 3-hydroxybutyrate, was found to be an endogenous ligand for HCA2 (Taggart et al., 2005). HCA2 is predominantly expressed in adipose tissue. Upon binding to niacin, HCA2 is coupled to the G protein and lowers cAMP levels in adipocytes through inhibition of adenylyl cyclase, resulting in decreased hormone-sensitive lipase activity and reduced hydrolysis of triglycerides to free fatty acids (Soga et al., 2003; Tunaru et al., 2003; Wise et al., 2003). Several studies have indicated that niacin-induced flushing is also mediated by HCA2 through the release of prostaglandin D2 and prostaglandin E2 (Benyó et al., 2005; Meyers et al., 2007). In addition, new nonflushing drugs with low toxicity based on the pharmacological effects of niacin that act via HCA2 are in development to treat dyslipidemia and prevent cardiovascular disease (Kamanna and Kashyap, 2007). Richman et al. (2007) reported that agonists that fail to significantly activate ERK1/2 phosphorylation and do not cause rapid receptor internalization do not lead to a flushing response. Therefore, an understanding of the regulation of HCA2 signaling and trafficking is important for the potential development of pharmaco- therapeutic agents with antilipolytic activity that are also nonflushing.

Our previous studies have demonstrated that the membrane-bound Gαi3 subunit is dissociated from Gt upon activation of HCA2 by niacin. This is followed by recruitment of the C-terminal tail in activating receptor desensitization, endocytosis, and internalization. We also attempted to identify structural motifs within the C-terminal tails of HCA receptors that are responsible for down- stream signaling pathway activation and direct interactions between the HCA2 receptor and the intracellular sorting machinery. We demonstrated for the first time that several distinct domains within the C-terminal tail of HCA2 are responsible for the regulation of HCA2 receptor export, phosphorylation, desensitization, and internalization.

Materials and Methods

Materials. Lipofectamine 2000 and G418 were purchased from Invitrogen (Carlsbad, CA). Cell culture media and fetal bovine serum were obtained from HyClone (Beijing, China). The pCMV-FLAG, pcDNA3.1, and pEGFP-N1 vectors were purchased from Sigma-Aldrich (St. Louis, MO), Invitrogen, and Clontech (Mountain View, CA), respectively. pDsRed2-ER was a gift from Dr. Tao Xu (Institute of Biophysics, Academy of Sciences, Beijing, China). 1,1-Dioctadecyl-3,3,3′,3′-tetramethylindocarbocyanine perchlorate (DiI) was purchased from Beyotime (Haimen, China). Monoclonal anti-FLAG M2 antibody, monoclonal anti-HA M1 antibody, and horseradish peroxidase (HRP)-conjugated anti-mouse IgG were obtained from Sigma-Aldrich. Anti-phospho-ERK1/2 (Thr202/Tyr204), ERK1/2 antibodies, and HRP-conjugated anti-rabbit IgG were obtained from Cell Signaling Technology (Danvers, MA).

Cell Culture and Transfection. HEK-293 cells were maintained in Dulbecco’s modified Eagle’s medium supplemented with 10% heat-inactivated fetal bovine serum and 4 mM L-glutamine. Plasmid constructs were transfected or cotransfected into HEK-293 cells using Lipofectamine 2000 according to the manufacturer’s instructions. All cells were incubated at 37°C in a humidified chamber with 5% CO2 and 95% air.

Molecular Cloning and Plasmid Construction. For fluorescence detection experiments, the cDNA of the HCA2 receptor was C-terminally fused to the cDNA of enhanced green fluorescent protein (EGFP). To perform enzyme-linked immunosorbent assays (ELISA) for quantification of receptor expression and internalization, the cDNA of the HCA2 receptor was inserted between the HindIII and EcoRI sites of the pCMV-FLAG vector as described previously (Li et al., 2010). Mutations and truncations were introduced by overlap extension polymerase chain reaction. All constructs were sequenced to verify sequences and orientations. The arrestin3-EGFP was generated as described previously (Li et al., 2011).

Measurement of Cell Surface and Total Cellular Receptor Levels by ELISA. To assess levels of cell surface receptors, cells transiently transfected with either wild-type (WT) or mutant FLAG-HCA2 cDNA constructs were fixed with 3.7% formaldehyde in Tris-buffered saline (TBS) for 5 min, washed three times with TBS, and then blocked for 1 h with 1% BSA/TBS. Cells were then incubated at room temperature with anti-FLAG M2 antibody (1:1000) diluted in TBS (1% BSA) for 1 h, probed with HRP-labeled secondary antibody, and chased in chemiluminescence using HRP substrate (Sigma-Aldrich). Development was stopped by removing 0.1 ml of the substrate to a 96-well microtiter plate containing 0.1 ml of 1% SDS. Plates were read at 405 nm in a microplate reader (Bio-Rad Laboratories, Hercules, CA) using Microplate Manager software.

Measurement of Cell Surface and Total Cellular Receptor Levels by ELISA. To assess levels of cell surface receptors, cells transiently transfected with either wild-type (WT) or mutant FLAG-HCA2 cDNA constructs were fixed with 3.7% formaldehyde in Tris-buffered saline (TBS) for 5 min, washed three times with TBS, and then blocked for 1 h with 1% BSA/TBS. Cells were then incubated at room temperature with anti-FLAG M2 antibody (1:1000) diluted in TBS (1% BSA) for 1 h, probed with HRP-labeled secondary antibody, and chased in chemiluminescence using HRP substrate (Sigma-Aldrich). Development was stopped by removing 0.1 ml of the substrate to a 96-well microtiter plate containing 0.1 ml of 1% SDS. Plates were read at 405 nm in a microplate reader (Bio-Rad Laboratories, Hercules, CA) using Microplate Manager software.

Quantitative Measurement of Cell Surface Receptor Internalization. Cell surface receptor internalization was quantitatively assessed by ELISA as described previously (Li et al., 2010).

Measurement of Receptor Localization and Internalization by Confocal Imaging. To measure receptor localization, HEK-293 cells were transiently transfected with WT or mutant HCA2-EGFP with and without pDsRed2-ER. On the following day, cells were seeded in coverage, glass-bottomed six-well plates and then fixed with 3% paraformaldehyde for 15 min. For the internalization assay, HEK-293 cells stably expressing WT or mutant HCA2-EGFP were treated with 200 μM niacin for 1 h. After removal of the agonist, cells were fixed with 3% paraformaldehyde for 15 min. Confocal images were taken on a Zeiss LSM 510 microscope with an attached Axiovert 200 microscope and LSM5 computer system.

Role of C Terminus in Hydroxycarboxylic Acid Receptor-2
**Immunofluorescence Microscopy.** HEK-293 cells were transfected with WT or mutant FLAG-HCA2. Then cells were incubated with anti-FLAG-FTTC antibody (1:250 dilution) for 1 h at 4°C in DMEM supplemented with 1% BSA. Cells were washed twice with PBS and then were incubated in media alone (vehicle) or with 200 μM niacin for 30 min at 37°C. Cells were then fixed with 3% paraformaldehyde-PBS for 15 min, and receptor-antibody complexes were visualized by confocal microscopy.

**cAMP Accumulation.** HEK-293 cells transiently cotransfected with HCA2 or HCA2 mutants and pCRE-Luc were grown to 90 to 95% confluence in DMEM without fetal bovine serum. Cells were then stimulated with 10 μM forskolin alone or 10 μM forskolin with different concentrations of niacin and incubated for 4 h at 37°C. Luciferase activity was detected using a firefly luciferase kit (Promega, Madison, WI). For the receptor desensitization assay, HEK-293 cells transiently transfected with HCA2 or HCA2 mutants were pretreated with water (control) or 200 μM niacin for 1 h, washed three times with ice-cold PBS, and stimulated with 10 μM forskolin and 200 μM niacin for 20 min. The reaction was stopped by a quick wash with ice-cold PBS followed by the addition of cell lysis buffer. The cAMP concentration was assessed using a commercially available cAMP detection kit (R&D Systems, Minneapolis, MN).

**Synthesis of siRNAs and siRNA Transfection.** All arrestins and clathrin siRNAs were chemically synthesized by Dharmacon RNA Technologies (Lafayette, CO). The transfection protocols for arrestin and clathrin siRNAs were reported previously (Li et al., 2010). Forty-eight hours after transfection, cells were split for the indicated assay on the following day.

**Immunoblotting and Immunoprecipitation.** Phosphorylated ERK immunoblotting using the antibody anti-phospho-ERK1/2 (diluted 1:2000; Cell Signaling Technology) was performed as described previously (Zhou et al., 2012). Total ERK1/2 was detected using anti-ERK1/2 (diluted 1:2000; Cell Signaling Technology). Clathrin was detected using mouse monoclonal antibody anti-clathrin HC (TD.1, diluted 1:2000; Santa Cruz Biotechnology, Inc., Santa Cruz, CA). For immunoprecipitation, cells were treated with serum-free media with or without niacin. Cells were washed once with PBS at 4°C, harvested by gentle scraping, pelleted, and resuspended in glycerol lysis buffer including complete protease inhibitor. Lysates were normalized for equal protein concentrations and immunoprecipitated with conjugated M2 beads (Sigma-Aldrich). Immunoprecipitation reactions were incubated at 4°C for 1 h, washed three times with glycerol lysis buffer, and resuspended in SDS running buffer. Samples were subjected to SDS-PAGE analysis and Western blotting with anti-clathrin HC (TD.1) antibody. The blots were stripped and reprobed using mouse monoclonal anti-FLAG antibody (1:1000) as a control for protein loading.

** Arrestin Translocation Assay.** The fluorescence detection experiment of arrestin translocation was performed as described previously (Li et al., 2011). The method of quantitative arrestin3 membrane translocation was described previously (Li et al., 2010).

**Analysis of Receptor Phosphorylation by Gel Shift Analysis.** Plasmid constructs of Flag-tagged HCA2 receptor wild-type or SSTA/A mutant were transiently transfected into HEK-293 cells and stably selected using G418. Cells grown in six-well plates were incubated in Dulbecco’s modified Eagle’s medium for 24 h at 37°C to reach ~90% confluence. Cells were stimulated with water (control) or 200 μM niacin for 10 min, washed twice with ice-cold PBS and then with 0.3 ml of ice-cold lysis buffer containing 150 mM NaCl, 20 mM Hepes, pH 7.2, 5 mM EDTA, and 1% Triton X-100, and one complete EDTA-free protease inhibitor cocktail tablet (Roche, Mannheim, Germany) was added. The plates were rocked for 30 min at 4°C, and supernatants were collected by centrifugation at 12,000g for 20 min at 4°C. The samples were subjected to electrophoresis on a 10% polyacrylamide gel for 2 h at 120 V and then transferred to nitrocellulose. After blocking with 5% milk, blots were probed for HCA2 receptor with 1:500 anti-Flag mouse antibody (M2 antibody; Sigma-Aldrich), HRP-conjugated anti-mouse secondary antibody, and chemiluminescent detection.

**Results**

**Expression and Functional Characterization of C-Terminal Deletion Mutants of Human HCA2.** Multiple domains and motifs within the C-terminal tail of GPCRs have been implicated in the regulation of receptor expression, dimerization, internalization, trafficking, and signaling (Pankevych et al., 2003; Levoye et al., 2006; Delhaye et al., 2007; Labasque et al., 2008; McCulloch et al., 2008). We constructed an expression vector containing the HCA2 receptor fused with EGFP at the carboxyl terminus for direct visualization of HCA2 receptor membrane localization and trafficking. To identify the functional domains within HCA2 that regulate receptor localization, internalization, desensitization, and trafficking, we constructed a series of C-terminal deletion mutants of human HCA2 (Fig. 1a). We first examined the functional signaling of HCA2-EGFP by assaying cAMP accumulation. As shown in Supplemental Fig. 1, treatment of HEK-293 cells that transiently transfected HCA2-EGFP with niacin inhibited forskolin-stimulated cAMP production comparably to that in cells expressing FLAG-HCA2. These results demonstrate that a fusion of the C terminus of HCA2 receptors to EGFP functions similarly to the untagged HCA2 receptor.

Confocal microscopy revealed that HCA2 and the Δ315–328 and Δ344–363 mutants were expressed and colocalized with the membrane marker DiI with minimal intracellular accumulation. In contrast, there was nearly no expression of the Δ295–314 construct, and only half of the expressed Δ329–343 construct colocalized with DiI (Fig. 1b). The same result was also observed by ELISA analysis (Fig. 1c). The decrease in cell surface expression was not simply attributable to deficient cellular expression of the mutant receptor proteins, because similar total cellular levels of HCA2 wild-type protein and all mutant proteins with C-terminal deletions were detected by ELISA analysis after cell permeabilization (Fig. 1d). These data suggest that the C-terminal tail of HCA2 is important for the correct localization of the HCA2 receptor in the plasma membrane.

We next examined whether signaling of the HCA2 C-terminal deletion mutants was functional by assaying CRE-driven luciferase activity and ERK1/2 activation. As shown in Fig. 1, e and f, the Δ315–328 and Δ343–363 mutants functioned identically to the intact HCA2 receptor. However, the Δ329–343 mutant showed decreased activity in the inhibition of forskolin-induced luciferase activity, whereas the Δ295–314 mutant exhibited complete loss in agonist-induced inhibition of forskolin-stimulated CRE-driven luciferase activity (Fig. 1e). Identical results were observed for ERK1/2 activation (Fig. 1f). Taken together, these data suggest that the C-terminal tail of HCA2 plays an important role in the regulation of receptor cell surface localization but not in the modulation of receptor expression and signaling.

**The C-Terminal Domain between Residues Tyr295 and Gln314 Plays an Essential Role in the Regulation of HCA2 Export from the Endoplasmic Reticulum to the Cell Surface.** To determine whether the stably expressed Δ295–314 mutant undergoes internalization upon agonist binding, HEK-293 cells stably transfected with...
HCA₂-EGFP or Δ295–314-EGFP were incubated with 200 μM niacin for 40 min at 37°C and examined by confocal microscopy. As shown in Fig. 2a, wild-type HCA₂ receptors were internalized from the cell surface into the cytoplasm with a punctate distribution upon activation by niacin, whereas the Δ295–314 mutant was still localized primarily in the cytoplasm.

We proceeded to further investigate the intracellular compartments into which the Δ295–314 mutant was retained. The wild-type and Δ295–314 mutant were coexpressed with
the endoplasmic reticulum (ER) marker DsRed2-ER, and their respective colocalization was analyzed by confocal microscopy. As anticipated, the wild type showed no colocalization with DsRed2-ER. In contrast, the \( \text{H9004}^{295–314} \) mutant was extensively colocalized with DsRed2-ER (Fig. 2b). These results are consistent with previous studies showing that the F(X)\(_6\)LL motif in the C termini of GPCRs is essential for the export of receptors from the ER to the cell surface (Duvernay et al., 2004, 2009) and suggesting that the F(X)\(_6\)LI motif may function as a signal dictating the ER export of HCA\(_2\).

The C-Terminal Ser/Thr cluster of 326STS328 Is Involved in the Regulation of Arrestin Recruitment, Receptor Internalization and Phosphorylation, and Signaling Desensitization of HCA\(_2\). We next used confocal imaging and ELISA analysis to assess the ability of HCA\(_2\) and HCA\(_3\) mutants with C-terminal deletions to undergo agonist-induced internalization in stably transfected HEK-293 cells. As shown in Fig. 3, a and b, after niacin stimulation (200 \( \mu \)M), both the \( \Delta 329–343 \) and \( \Delta 344–363 \) mutants displayed agonist-induced internalization comparable to that of the wild-type receptor, whereas the \( \Delta 315–328 \) mutant exhibited a significant decrease in niacin-mediated internalization. The domain between residues 315 and 328 seems to play a critical role in HCA\(_2\) internalization. These results, together with the potent inhibition of receptor internalization caused by GRK2 knockdown (Li et al., 2010), suggest that the serine and threonine residues within this domain may serve as GRK phosphorylation sites and may be critical for arrestin binding and agonist-induced receptor internalization. To address the potential functional role of the serine and threonine residues within this domain, we constructed site-directed mutants that substituted alanines for the serine and threonine residues within this domain. These mutants were then stably transfected into HEK-293 cells. The replacement of threonine 318 with alanine resulted in a loss of inhibition of niacin-induced internalization compared with that for wild-type HCA\(_2\). However, the niacin-induced loss of cell surface receptors in HEK-293 cells expressing the S326A, T327A, or S328A mutants was significantly decreased, and the STS/A mutant exhibited an \( \sim 75\% \) decrease in receptor internalization compared with that for wild-type HCA\(_2\). Although the S326A, T327A, S328A, and STS/A mutants were resistant to agonist-induced rapid internalization, there were no differences in CRE-driven luciferase activity or niacin-induced ERK1/2 phosphorylation between these mutants and the wild-type HCA\(_2\) (Fig. 3, e and f).

Our previous studies have demonstrated that niacin-activated HCA\(_2\) is phosphorylated by GRK2 followed by binding of arrestin3 at the C-terminal tail, which targets HCA\(_2\) receptors to clathrin-coated pits and causes their internalization in HEK-293-HCA\(_2\) cells (Li et al., 2010). To further investigate the interaction of these HCA\(_2\) mutants with ar-
restin3, we cotransfected HCA2 wild-type, Δ315–328, STS/A, and Δ329–343 stably expressing HEK-293 cells with arrestin3-EGFP and visualized the translocation of the latter after agonist-mediated receptor activation. In the absence of niacin, arrestin3 was localized primarily in the cytoplasm in both the wild-type and mutant cells (Fig. 4a). Stimulation with niacin resulted in robust recruitment of arrestin3-GFP to the cell membrane in the wild-type HCA2-expressing cells (Fig. 4a), but no redistribution of arrestin3-GFP was observed upon agonist activation in the Δ315–328, STS/A, and Δ329–343 mutant-expressing cells.

Thus, the STS/A mutant exhibited a significant decrease in niacin-mediated internalization and arrestin3 translocation. These results, together with the potent inhibition of receptor internalization caused by GRK2 knockdown (Li et al., 2010), suggest that the serine and threonine residues within this domain may serve as GRK2 phosphorylation sites. To address this hypothesis, we transfected HEK-293 cells with
either HCA2 wild-type or STS/A mutant receptor and analyzed receptor phosphorylation by gel shift analysis. As shown in Fig. 4b, in the absence of niacin, both wild-type and STS/A mutant HCA2 receptors migrate as a major band of ~43 kDa. Treatment of cells expressing the wild-type receptor with 200 μM niacin resulted in a significant mobility shift of the receptor with a few slower migrating bands being observed. In striking contrast, this agonist-promoted shift was not observed in cells expressing the STS/A mutant HCA2 receptor. We attribute this mobility shift to agonist-promoted phosphorylation of the receptor, which is lost in the STS/A mutant.

Next, we further evaluated the agonist-induced desensitization of these HCA2 mutants. Cells were pretreated with vehicle (control) or with 200 μM niacin for 1 h, washed, and incubated with 200 μM niacin and 10 μM forskolin for 20 min. cAMP accumulation was then determined using a commercially available kit. Compared with wild-type HCA2, the Δ315–328 and STS/A mutants were impaired in agonist-induced rapid receptor desensitization (Fig. 4c). Taken together, our findings strongly suggest that the serine/threonine cluster of residues from 326 to 328 plays an essential role in receptor internalization and phosphorylation, desensitization, and association with arrestin3.
Deletion of the C-Terminal Domain from Val329 to Glu343 Leads to Constitutive Activation of HCA2. As shown above, the data obtained from confocal microscopy and ELISA analysis showed that only half of the expressed H9004329–343 mutant protein localized to the plasma membrane. To determine the intracellular compartments to which the intracellular H9004329–343 mutant was localized, the H9004329–343 mutant was coexpressed with the ER marker DsRed2-ER, and their colocalization was analyzed. Of interest, the H9004329–343 mutant receptor did not significantly colocalize with DsRed2-ER in the cytoplasm (Fig. 5a). To evaluate the primary source of intracellular H9004329–343 receptors, the distribution of H9004329–343-EGFP was monitored after treatment (1, 2, and 4 h) with cycloheximide (CHX; 25 μM). HEK-293 cells stably expressing H9004329–343-EGFP were incubated with either 100 μg/ml Alexa Fluor 546-labeled transferrin (red) or 50 nM LysoTracker DND-99 (red) for 40 min. All images are representative of at least three independent experiments. CTL, control.

Fig. 5. Cytoplasmic localization of the Δ329–343 mutant receptor. a, HEK-293 cells were transiently cotransfected with HCA2-EGFP or Δ329–343-EGFP (green) and pDsRed2-ER (red). After 24 h, cells were fixed and examined by confocal microscopy. b, HEK-293 cells stably expressing Δ329–343-EGFP were treated (1, 2, and 4 h) with cycloheximide (CHX; 25 μM). c, HEK-293 cells stably expressing Δ329–343-EGFP (green) were incubated with either 100 μg/ml Alexa Fluor 546-labeled transferrin (red) or 50 nM LysoTracker DND-99 (red) for 40 min. All images are representative of at least three independent experiments. CTL, control.
bated at 4°C with monoclonal anti-FLAG-FITC (M2) antibody (1:250) for 1 h in DMEM with 1% BSA. After washing three times with PBS, cells were incubated in media alone (vehicle) or with 200 μM niacin for 30 min at 37°C. Receptor-antibody complexes were then analyzed by confocal microscopy (a) or ELISA (b). c and d, HEK-293 cells stably expressing Δ329–343 were incubated in media alone (vehicle) or with cycloheximide (CHX) or with cycloheximide and MβCD (5 mM) or sucrose (450 mM) for 2 h and then were analyzed by confocal microscopy (c) or ELISA (d). e and f, arrestin and clathrin expression was measured as described under Materials and Methods. g, The role of arrestin2/3 and clathrin in the regulation of Δ329–343 constitutive endocytosis was measured by confocal microscopy. h, HEK-293 cells transiently cotransfected with HCA2 or Δ329–343 mutant and pCRE-Luc were stimulated with media alone or with 10 μM forskolin for 4 h at 37°C. Luciferase activity was detected by a firefly luciferase kit. i, HEK-293 cells stably expressing wild-type HCA2 or Δ329–343 mutant were stimulated with media alone or with 10 μM forskolin for 20 min. The cAMP concentration was assessed with a commercially available kit. Data were analyzed by using Student’s t test (*, p < 0.05; **, p < 0.01; ***, p < 0.001). All images and data shown are representative of at least three independent experiments. IB, immunoblot; CTL, control.
and 343 within the C-terminal tail of HCA2 probably plays a significant role in regulating HCA2 internalization and desensitization. Our findings suggest that a sequence between residues 329 and 343 within the C-terminal tail of HCA2 is effective and specifically knocked down by the specific siRNA treatment and was unaffected in cells treated with a nonspecific or control siRNA (Fig. 6, e and f). Silencing clathrin but not arrestin2/3 effectively inhibited ligand-independent endocytosis of Δ329–343 as analyzed by microscopy (Fig. 6g). Taken together, these data suggest that Δ329–343 undergoes constitutive endocytosis in a clathrin-dependent and arrestin2/3-independent pathway.

To further evaluate whether the Δ329–343 mutant is constitutively active in receptor signaling, we examined the ligand-independent levels of cAMP in cells expressing the mutant or the wild-type receptor. HEK-293 cells transiently cotransfected with the HCA2 wild-type or Δ329–343 mutant receptor and pCRE-Luc were grown to 90 to 95% confluence and stimulated with dimethyl sulfoxide alone or 10 μM forskolin in DMEM without fetal bovine serum. After incubation for 4 h at 37°C, luciferase activity was detected. We found that expression of the Δ329–343 mutant resulted in significantly lower basal and forskolin-stimulated cAMP levels (P < 0.01) than expression of the wild-type receptor (Fig. 6h). This result was confirmed using a nonradioactive competitive binding cAMP kit (Fig. 6i), indicating that the Δ329–343 mutant is constitutively active in Gs-coupled signaling. Our findings suggest that a sequence between residues 329 and 343 within the C-terminal tail of HCA2 probably plays a crucial role in keeping HCA2 in an inactive conformation.

Discussion

The rapid internalization or sequestration of receptors into intracellular membrane compartments after agonist activation is important for the regulation of GPCR signaling. Our previous results indicated the involvement of GRK2 and arrestin3 in the regulation of HCA2 receptor internalization. Intracellular GPCR domains, particularly the C terminus, are generally thought to regulate receptor desensitization, internalization, and intracellular trafficking by facilitating association of the receptor with various cellular proteins. To understand more about the structural requirements for the regulation of HCA2 internalization and desensitization, we first constructed four mutants, each with a deletion of 10 to 39 amino acids at the C-terminal tail, which were distinct from progressive truncated mutants. We used these mutants to successfully identify different domains responsible for receptor export, constitutive activity, desensitization, and internalization. A comprehensive series of mutants with alanine substitutions replacing conserved serine and threonine residues within the C-terminal tail was also generated to pinpoint the key residues essential for GRK2-mediated phosphorylation and arrestin3 association.

A regulatory mechanism shared by most GPCRs involves GRK-mediated phosphorylation of the C terminus of the receptor followed by arrestin binding, which leads to receptor internalization (Lefkowitz and Shenoy, 2005). Prior studies have examined the role of the carboxyl-terminal tail and putative domains for GRK phosphorylation in the regulation of GPCR internalization. A previous study has demonstrated that a cluster of serine/threonine residues in the C-terminal tail is essential for receptor internalization and desensitization, serving as targets for agonist-induced phosphorylation and association of the arrestin adaptor proteins (Oakley et al., 2001). A bradykinin B2 receptor mutant with four alanine substitutions in a serine/threonine cluster in the C-terminal tail has also been shown to be deficient in arrestin binding and in receptor internalization (Zimmerman et al., 2011). Furthermore, substitution of serine 355, 356, and 364 of the β2-adrenergic receptor to alanines reduces GRK-mediated C-terminal phosphorylation by at least 90% and almost completely eliminates receptor internalization (Seibold et al., 2000). In the current study, using a series of mutants with C-terminal deletions of 15 to 20 amino acids out of a total of 69 residues, we showed that the sequence from Arg315 to Ser328 (Δ315–328) is required for agonist-mediated receptor phosphorylation and internalization. Further mutagenesis studies using alanine substitution defined the three residues (Ser326, Thr327, and Ser328) responsible for GRK-mediated phosphorylation and internalization. We previously showed that HCA2 internalization can be triggered by arrestin3, but not by arrestin2, using siRNA-mediated knockdown (Li et al., 2010). Our findings lead to the conclusion that, upon activation by niacin, GRK2 mediates the phosphorylation of Ser326, Thr327, and Ser328 residues in the carboxyl tail of HCA2. This then facilitates arrestin3 binding to the receptor, resulting in receptor internalization by targeting receptors to clathrin-coated pits. The recruitment of arrestins to phosphorylated receptors results in attenuation of the GPCR response by uncoupling receptors from heterotrimeric G proteins. This mechanism of GPCR desensitization is important in allowing cells to avoid agonist overstimulation. It is generally accepted that both the second messenger-dependent protein kinases, PKA and PKC, and GRKs are involved in the phosphorylation of serine and threonine residues within the intracellular loops and C-terminal tail domains of GPCRs (Ferguson, 2001). For GPCRs, phosphorylation of the PKA and PKC sites within the third intracellular loop and the carboxyl-terminal tail has been found to contribute to agonist-dependent desensitization (Yuan et al., 1994; Moffett et al., 1996; Liang et al., 1998; Tang et al., 1998), whereas mutation of all of the serine and threonine residues within the C-terminal tail of the β2-adrenergic receptor and the third intracellular loop of the m2 muscarinic acetylcholine receptor prevents the induction of GRK-mediated desensitization (Bouvier et al., 1988; Nakata et al., 1994). Results from our previous mutagenesis experiments demonstrated that a sequence of 14 amino acids, from residues 315 to 328 (Δ315–328) at the C-terminal end of the HCA2 receptor is responsible for GRK-mediated phosphorylation and arrestin recruitment and internalization (Li et al., 2011). In the present study, mutant receptors with C-terminal deletions between Arg315 to Ser328 (Δ315–328) or with alanine substitutions for the three Ser/Thr residues in this region (Ser326, Thr327, and Ser328) not only exhibited deficiencies in arrestin3 binding and receptor internalization and phosphorylation (Figs. 3, c and d, and 4, a and b) but also exhibited reduced agonist-
induced desensitization during the inhibition of forskolin-induced cAMP accumulation (Fig. 4c). These results are consistent with our previous observation that the Δ315–328 mutant exhibited sustained activation of ERK1/2 from 10 to 30 min compared with wild-type HCA2 (Li et al., 2011). From our current data, we cannot exclude a potential role for PKA and PKC in regulating HCA2 desensitization. However, GRK-mediated phosphorylation of Ser326, Thr327, and Ser328 is also likely to contribute to the agonist-dependent desensitization of the HCA2 receptor.

Another interesting observation of the current study was that mutant HCA2 receptors with a 15-amino acid C-terminal deletion from Val329 to Glu343 showed significant constitutive endocytosis (Fig. 6, a and b) compared with wild-type HCA2. Moreover, this ligand-independent internalization of the Δ329–343 mutant underwent a clathrin-dependent and arrestin2/3-independent pathway (Fig. 6g). It is now well established that GPCRs can be activated spontaneously or constitutively in the absence of an agonist. The first observation of constitutive GPCR activity was obtained for the δ-opioid receptor (Koski et al., 1982) and the β2-adrenergic receptor (Cerione et al., 1984). During the past decades, more than 60 GPCRs have been shown to be constitutively active in both recombinant cells and native cells (Costa and Herz, 1989; Cotechia et al., 1990; Smit et al., 2007). In addition, naturally occurring mutant GPCRs with constitutive activities greater than wild-type activity are responsible for a variety of human diseases (Seifert and Wenzel-Seifert, 2002). Considerable evidence has accumulated in recent years to suggest that specific sequences in the C-terminal regions of GPCRs play a role in mediation of constitutive receptor activity. Deletion of the last several residues of the thyrotropin-releasing hormone receptor and the prostaglandin E receptor results in constitutive receptor activity (Matus-Leibovich et al., 1995; Hasegawa et al., 1996), and shorter C-terminal variants of 5-HT1 receptors have also been shown to have constitutive activity (Clayesen et al., 1999). In addition, deletion of the C-terminal regions of the β-adrenergic receptor (Parker and Ross, 1991) and the CB1 cannabinoid receptor (Nie and Lewis, 2001) promotes their constitutive activity. Mutation of the proximal palmitoylation site C404S of the 5-HT7(a) receptor leads to ligand-independent internalization of the Δ329–343 mutant underwar a clathrin-dependent and arrestin2/3-independent pathway (Fig. 6g). It is now well established that GPCRs can be activated spontaneously or constitutively in the absence of an agonist. The first observation of constitutive GPCR activity was obtained for the δ-opioid receptor (Koski et al., 1982) and the β2-adrenergic receptor (Cerione et al., 1984). During the past decades, more than 60 GPCRs have been shown to be constitutively active in both recombinant cells and native cells (Costa and Herz, 1989; Cotechia et al., 1990; Smit et al., 2007). In addition, naturally occurring mutant GPCRs with constitutive activities greater than wild-type activity are responsible for a variety of human diseases (Seifert and Wenzel-Seifert, 2002). Considerable evidence has accumulated in recent years to suggest that specific sequences in the C-terminal regions of GPCRs play a role in mediation of constitutive receptor activity. Deletion of the last several residues of the thyrotropin-releasing hormone receptor and the prostaglandin E receptor results in constitutive receptor activity (Matus-Leibovich et al., 1995; Hasegawa et al., 1996), and shorter C-terminal variants of 5-HT1 receptors have also been shown to have constitutive activity (Clayesen et al., 1999). In addition, deletion of the C-terminal regions of the β-adrenergic receptor (Parker and Ross, 1991) and the CB1 cannabinoid receptor (Nie and Lewis, 2001) promotes their constitutive activity. Mutation of the proximal palmitoylation site C404S of the 5-HT7(a) receptor leads to ligand-independent internalization of the Δ329–343 mutant underwar a clathrin-dependent and arrestin2/3-independent pathway (Fig. 6g).

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Authorship Contributions

Participated in research design: Li, Zhou, J.Lu, and Benovic.
Conducted experiments: Li, Zhou, Yu, Shi, and J.Luo.
Contributed new reagents or analytic tools: Li, Zhou, and J.Luo.
Performed data analysis: Li, Zhou, Yu, Chen, and Shi.
Wrote or contributed to the writing of the manuscript: Zhou, J.Lu, Benovic, and Li.

References


Hasegawa H, Negishi M, and Ichikawa A (1996) Two isoforms of the human HCA2 receptor also seem likely to play a critical role in receptor desensitization from the ER to the cell surface.

We conclude that several distinct C-terminal domains within the C-terminal tail of the human HCA2 receptor directly link this receptor to interaction with the intracellular sorting machinery and to the phosphorylation, internalization, and desensitization pathways. Our findings provide a structural and mechanistic basis for better understanding the regulation of desensitization and internalization of GPCRs.


